CETIFICATION

SDG No:

JC16038

Laboratory:

Accutest, New Jersey

Site:

BMSMC, Building 5 Area

Matrix:

Groundwater

SM04.00.06 Humacao, PR

SUMMARY:

Groundwater samples (Table 1) were collected on the BMSMC facility – Building 5 area. The BMSMC facility is located in Humacao, PR. Samples were taken March 8-9, 2016 and were analyzed in Accutest Laboratory of Dayton, New Jersey that reported the data under SDG No.: JC16038. Results were validated using the latest guidelines (July, 2015) of the EPA Hazardous Waste Support Section and the QC criteria for SW 846 methods, latest revision, for low molecular weight alcohols (LMWA). The analyses performed are shown in Table 1. Individual data review worksheets are enclosed for each target analyte group. Data sample organic data samples summary form shows for analytes results that were qualified.

In summary the results are valid and can be used for decision taking purposes.

Table 1. Samples analyzed and analysis performed

SAMPLE ID	SAMPLE	ANALYSIS PERFORMED
	DESCRIPTION	
JC16038-1	MW-13	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-2	MW-7	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-3	MW-3	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-4	MW-5	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-5	MW-16	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-6	MW-16	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-7	TB030902	VOCs; LMWA
JC16038-8	S-30	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-9	FB030816	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA
JC16038-10	MW-11	VOCs; SVOCs; NAPHTHALENE; 1,4-DIOXANE (SIM);
		PESTICIDES; LMWA

Reviewer Name:

Rafael Infante

Chemist License 1888

Signature:

Date:

April 16, 2016

Page 1 of 2

Client Sample ID: MW-13

Lab Sample ID: JC16038-1

Matrix: Method: AO - Ground Water

SW846 8260C

Date Received: 03/11/16

Date Sampled: 03/09/16

Percent Solids: n/a

Project: BMSMC, Building 5 Area, PR

File ID Run #1 U204171.D DF Analyzed 1 03/16/16

By NH Prep Date n/a

Prep Batch n/a

Q

J

j

Analytical Batch VU9384

Run #2

Purge Volume

Run #1 5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Unit
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93- 3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	0.28	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	0.90	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	ND	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	ND	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	ND	1.0	0.27	ug/l
76-13-1	Freon 113	30.1	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l
					43

Rafael Infant Méndez LIC # 188

ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 2

Client Sample ID: MW-13

Lab Sample ID: JC16038-1

Matrix: Method:

AQ - Ground Water SW846 8260C

Project:

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	ND	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	ND	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	ND	1.0	0.38	ug/l	
95-47-6	o-Xylene	ND	1.0	0.17	ug/l	
1330-20-7	Xylene (total)	ND	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limi	ts	
1868-53-7	Dibromofluoromethane	106%		76-12	20%	
17060-07-0	1,2-Dichloroethane-D4	108%		73-12	22%	
2037-26-5	Toluene-D8	99%		84-13	19%	
460-00-4	4-Bromofluorobenzene	103%		78-11	17%	



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B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

By

SD

Client Sample ID: MW-13

Lab Sample ID:

JC16038-1

AQ - Ground Water

DF

1

Date Sampled: 03/09/16

Matrix: Method:

SW846 8270D SW846 3510C

Date Received: 03/11/16 Percent Solids: n/a

Project:

BMSMC, Building 5 Area, PR

Prep Batch

Analytical Batch

Run #1 Run #2

F155577.D

File ID

Final Volume

Analyzed

03/16/16

OP92078

Q

Prep Date

03/15/16

EF6543

Initial Volume 1000 ml 1.0 ml

Run #1 Run #2

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Units
95-57-8	2-Chlorophenol	ND	5.0	0.93	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.0	1.4	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.0	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.0	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	10	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.0	0.87	ug/l
95-48-7	2-Methylphenol	ND	2.0	0.82	ug/l
	3&4-Methylphenol	ND	2.0	0.67	ug/l
88-75-5	2-Nitrophenol	ND	5.0	1.4	ug/l
100-02-7	4-Nitrophenol	ND	10	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.0	1.4	ug/i
108-95-2	Phenol	ND	2.0	0.31	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.0	1.4	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.0	1.5	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.0	1.4	ug/l
83-32-9	Acenaphthene	ND	1.0	0.29	ug/l
208-96-8	Acenaphthylene	ND	1.0	0.24	ug/l
98-86-2	Acetophenone	ND	2.0	0.28	ug/l
120-12-7	Anthracene	ND	1.0	0.25	ug/l
1912-24-9	Atrazine	ND	2.0	0.42	ug/l
100-52-7	Benzaldehyde	ND	5.0	0.34	ug/l
56-55-3	Benzo(a)anthracene	ND	1.0	0.32	ug/l
50-32-8	Benzo(a)pyrene	ND	1.0	0.33	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.0	0.32	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.0	0.41	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.0	0.37	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.0	0.37	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.0	0.27	ug/l
92-52-4	1,1'-Biphenyl	ND	1.0	0.26	ug/l
91-58-7	2-Chloronaphthalene	ND	2.0	0.30	ug/l
106-47-8	4-Chloroaniline	1,3	5.0	0.23	ug/l
86-74-8	Carbazole	ND	1.0	0.29	ug/l



ND = Not detected

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Page 2 of 3

Client Sample ID: MW-13

Lab Sample ID: JC16038-1

Matrix: Method:

Project:

AO - Ground Water

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

ABN TCL List (SOM0 1.1)

11						
CAS No.	Compound	Result	RL	MDL	Units	Q
105-60-2	Caprolactam	ND	2.0	0.43	ug/l	
218-01-9	Chrysene	ND	1.0	0.35	ug/l	
111-91-1	bis(2-Chloroethoxy)methane	ND	2.0	0.26	ug/l	
111-44-4	bis(2-Chloroethyl)ether	ND	2.0	D.34	ug/l	
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.0	0.28	ug/l	
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.0	0.27	ug/l	
121-14-2	2,4-Dinitrotoluene	ND	1.0	0.26	ug/l	
606-20-2	2,6-Dinitrotoluene	ND	1.0	0.32	ug/l	
91-94-1	3,3'-Dichlorobenzidine	ND	2.0	0.53	ug/l	
53-70-3	Dibenzo (a,h) anthracene	ND	1.0	0.37	ug/l	
132-64-9	Dibenzofuran	ND	5.0	0.27	ug/l	
84-74-2	Di-n-butyl phthalate	ND	2.0	0.79	ug/l	
117-84-0	Di-n-octyl phthalate	ND	2.0	0.29	ug/l	
84-66-2	Diethyl phthalate	ND	2.0	0.24	ug/l	
131-11-3	Dimethyl phthalate	ND	2.0	0.31	ug/l	
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.0	0.77	ug/I	
206-44-0	Fluoranthene	ND	1.0	0.23	ug/l	
86-73-7	Fluorene	ND	1.0	0.29	ug/l	
118-74-1	Hexachlorobenzene	ND	1.0	0.42	ug/l	
87-68-3	Hexachlorobutadiene	ND	1.0	0.36	ug/l	
77-47-4	Hexachlorocyclopentadiene	ND	10	0.29	ug/l	
67-72-1	Hexachloroethane	ND	2.0	0.22	ug/l	
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.0	0.38	ug/l	
78-59-1	Isophorone	ND	2.0	0.29	ug/l	
91-57-6	2-Methylnaphthalene	ND	1.0	0.29	ug/l	
88-74-4	2-Nitroaniline	ND	5.0	0.21	ug/l	
99-09-2	3-Nitroaniline	ND	5.0	0.24	ug/l	
100-01-6	4-Nitroaniline	ND	5.0	0.34	ug/l	
91-20-3	Naphthalene	ND	1.0	0.28	ug/l	
98-95-3	Nitrobenzene	ND	2.0	0.46	ug/l	SOUCH
621-64-7	N-Nitroso-di-n-propylamine	ND	2.0	0.31	ug/l	386
86-30-6	N-Nitrosodiphenylamine	ND	5.0	0.29	ug/l	
85-01-8	Phenanthrene	ND	1.0	0.23	ug/l	Rotael I
129-00-0	Pyrene	ND	1.0	0.34	ug/l	Rofael I Mén
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.0	0.36	ug/l	: \\ 1.IC #
			757			10
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	Calinico Lic
367-12-4	2-Fluorophenol	42%		14-8	8%	S 11
4165-62-2	Phenol-d5	30%			10%	
	# nnvener6 1864			10-1	1070	



MDL = Method Detection Limit





RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Page 3 of 3

Client Sample ID: MW-13

Lab Sample ID: JC16038-1

Matrix:

AQ - Ground Water

Date Sampled: 03/09/16 Date Received: 03/11/16

Method: Project:

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Percent Solids: n/a

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limits
118-79-6	2,4,6-Tribromophenol	78%		39-149%
4165-60-0	Nitrobenzene-d5	75%		32-128%
321-60-8	2-Fluorobiphenyl	68%		35-119%
1718-51-0	Terphenyl-d14	73%		10-126%





MDL = Method Detection Limit

RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

E = Indicates value exceeds calibration range

Page 1 of 1

Client Sample ID: MW-13

Lab Sample ID: JC16038-1 Matrix:

AO - Ground Water

SW846 8270D BY SIM SW846 3510C

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

Project: BMSMC, Building 5 Area, PR

File ID DF Analyzed By Prep Date Prep Batch **Analytical Batch** Run #1 4P15636.D 1 03/17/16 IJ 03/15/16 OP92078A E4P804

Run #2

Method:

Initial Volume **Final Volume** Run #1 1000 ml $1.0 \, ml$

Run #2

CAS No. Compound Result RL MDL Units Q 91-20-3 Naphthalene ND 0.10 0.013 ug/l 123-91-1 1,4-Dioxane ND 0.10 0.053 ug/l CAS No. Surrogate Recoveries Run#1 Run# 2 Limits 4165-60-0 Nitrobenzene-d5 77% 24-125% 321-60-8 2-Fluorobiphenyl 77% 19-127% 1718-51-0 Terphenyl-d14 77% 10-119%



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

Page 1 of 1

Analytical Batch

GGH5213

Client Sample ID: MW-13

Lab Sample ID: JC16038-1

Matrix: Method:

Project:

AQ - Ground Water

SW846-8015C (DAI) BMSMC, Building 5 Area, PR Date Sampled: 03/09/16 Date Received:

03/11/16

Percent Solids: n/a

File ID DF Analyzed By Prep Date Prep Batch GH103819.D Run #1 XPL 1 03/18/16 n/a n/a

Run #2

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units	Q
64-17-5	Ethanol	ND	100	55	ug/l	
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l	
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l	
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l	
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l	
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l	
67-56-1	Methanol	ND	200	71	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
111-27-3	Hexanol	77%		56-1	45%	
111-27-3	Hexanol	75%		56-1	45%	





MDL = Method Detection Limit

N = Indicates presumptive evidence of a compound



RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

SGS Accutest

Report of Analysis

By

NH

n/a

Page 1 of 2

Client Sample ID: MW-7

Lab Sample ID: JC16038-2

File ID

U204139.D

Matrix: Method:

Project:

AQ - Ground Water

DF

1

SW846 8260C

BMSMC, Building 5 Area, PR

Analyzed

03/15/16

Date Sampled: 03/09/16

n/a

Q

J

Date Received: 03/11/16 Percent Solids: n/a

Prep Date Prep Batch **Analytical Batch**

VU9383

Run #1 Run #2

Purge Volume

Run #1 5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Unit
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	1.4	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/I
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	7:3	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	0.30	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	1:1	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	1.4	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	1.7	1.0	0.27	ug/l
76-13-1	Freon 113	ND	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l
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J = Indicates an estimated value ICEN

ND = Not detected MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

B = Indicates analyte found in associated method blank

fael Infante Méndez IC # 1881



Page 2 of 2

Client Sample ID: MW-7

Lab Sample ID: JC16038-2

Matrix: Method:

Project:

AQ - Ground Water SW846 8260C

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a



VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	ND	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	0.56	1.0	0.24	ug/l	J
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	0.66	1.0	0.22	ug/l	J
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	5.8	1.0	0.38	ug/l	
95-47-6	o-Xylene	0.50	1.0	0.17	ug/l	J
1330-20-7	Xylene (total)	6.3	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its	
1868-53-7	Dibromofluoromethane	105%		76-1	20%	
17060-07-0	1,2-Dichloroethane-D4	107%		73-1	22%	
2037-26-5	Toluene-D8	100%		84-1	19%	
460-00-4	4-Bromofluorobenzene	103%		78-1	17%	



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank N = Indicates presumptive evidence of a compound

Ву

SD

Client Sample ID: MW-7

Lab Sample ID: JC16038-2

File ID

Matrix: Method:

AO - Ground Water SW846 8270D SW846 3510C

Date Sampled: 03/09/16 Date Received: 03/11/16

Prep Date

03/15/16

Percent Solids: n/a

Project:

BMSMC, Building 5 Area, PR

Run #1

DF Analyzed F155578.D 1 03/16/16

Prep Batch

Q

Analytical Batch

OP92078 EF6543

Run #2

Initial Volume **Final Volume**

Run #1 950 ml

Run #2

1.0 ml

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Unite
95-57-8	2-Chlorophenol	ND	5.3	0.98	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.3	1.5	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.1	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.3	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	11	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.3	0.92	ug/l
95-48-7	2-Methylphenol	ND	2.1	0.86	ug/l
	3&4-Methylphenol	ND	2.1	0.71	ug/l
88-75-5	2-Nitrophenol	ND	5.3	1.5	ug/l
100-02-7	4-Nitrophenol	ND	11	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.3	1.5	ug/l
108-95-2	Phenol	ND	2.1	0.33	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.3	1.5	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.3	1.6	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.3	1.5	ug/l
83-32-9	Acenaphthene	ND	1.1	0.30	ug/l
208-96-8	Acenaphthylene	ND	1.1	0.25	ug/l
98-86-2	Acetophenone	ND	2.1	0.29	ug/l
120-12-7	Anthracene	ND	1.1	0.26	ug/l
1912-24-9	Atrazine	ND	2.1	0.44	ug/l
100-52-7	Benzaldehyde	ND	5.3	0.35	ug/l
56-55-3	Benzo(a)anthracene	ND	1.1	0.33	ug/l
50-32-8	Benzo(a)pyrene	ND	1.1	0.35	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.1	0.33	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.1	0.43	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.1	0.39	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.1	0.39	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.1	0.29	ug/l
92-52-4	1,1'-Biphenyl	ND	1.1	0.27	ug/l
91-58-7	2-Chloronaphthalene	ND	2.1	0.31	ug/l
106-47-8	4-Chloroaniline	ND	5.3	0.24	ug/l
86-74-8	· Caro continue			U.LT	ug/ i



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 3

Client Sample ID: MW-7

Lab Sample ID: JC16038-2

Matrix: Method:

Project:

AQ - Ground Water

SW846 8270D SW846 3510C

BMSMC, Building 5 Area, PR

Date Sampled: Date Received:

Q

03/09/16 03/11/16

Percent Solids: n/a

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Units
105-60-2	Caprolactam	ND	2.1	0.45	ug/l
218-01-9	Chrysene	ND	1,1	0.36	ug/l
111-91-1	bis(2-Chloroethoxy)methane	ND	2.1	0.27	ug/l
111-44-4	bis(2-Chloroethyl)ether	ND	2.1	0.36	ug/l
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.1	0.30	ug/l
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.1	0.29	ug/l
121-14-2	2,4-Dinitrotoluene	ND	1.1	0.28	ug/l
606-20-2	2,6-Dinitrotoluene	ND	1.1	0.34	ug/l
91-94-1	3,3'-Dichlorobenzidine	ND	2.1	0.56	ug/l
53-70-3	Dibenzo(a,h)anthracene	ND	1.1	0.38	ug/I
132-64-9	Dibenzofuran	ND	5.3	0.29	ug/l
84-74-2	Di-n-butyl phthalate	ND	2.1	0.83	ug/l
117-84-0	Di-n-octyl phthalate	ND	2.1	0.30	ug/l
84-66-2	Diethyl phthalate	ND	2.1	0.26	ug/l
131-11-3	Dimethyl phthalate	ND	2.1	0.33	ug/l
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.1	0.81	ug/l
206-44-0	Fluoranthene	ND	1.1	0.24	ug/l
86-73-7	Fluorene	ND	1.1	0.31	ug/l
118-74-1	Hexachlorobenzene	ND	1.1	0.44	ug/l
87-68-3	Hexachlorobutadiene	ND	1.1	0.38	ug/l
77-47-4	Hexachlorocyclopentadiene	ND	11	0.31	ug/l
67-72-1	Hexachloroethane	ND	2.1	0.23	ug/l
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.1	0.40	ug/l
78-59-1	Isophorone	ND	2.1	0.30	ug/l
91-57-6	2-Methylnaphthalene	ND	1.1	0.30	ug/l
88-74-4	2-Nitroaniline	ND	5.3	0.22	ug/l
99-09-2	3-Nitroaniline	ND	5.3	0.25	ug/l
100-01-6	4-Nitroaniline	ND	5.3	0.36	ug/l
91-20-3	Naphthalene	ND	1.1	0.30	ug/l
98-95-3	Nitrobenzene	ND	2.1	0.49	ug/l
621-64-7	N-Nitroso-di-n-propylamine	ND	2.1	0.33	ug/l
86-30-6	N-Nitrosodiphenylamine	ND	5.3	0.31	ug/l
85-01-8	Phenanthrene	ND	1.1	0.24	ug/l
129-00-0	Pyrene	ND	1.1	0.35	ug/l
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.1	0.38	ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its
367-12-4	2-Fluorophenol	44%		14-8	8%
4165-62-2	Phenol-d5	33%		10-1	10%



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 3 of 3

Client Sample ID: MW-7

Lab Sample ID: JC16038-2

Matrix: Method:

Project:

AQ - Ground Water

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Date Sampled: Date Received:

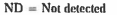
03/09/16 03/11/16

Percent Solids: n/a

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limits
118-79-6	2,4,6-Tribromophenol	85%		39-149%
4165-60-0	Nitrobenzene-d5	73%		32-128%
321-60-8	2-Fluorobiphenyl	68%		35-119%
1718-51-0	Terphenyl-d14	76%		10-126%





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J'= Indicates an estimated value

B = Indicates analyte found in associated method blank N = Indicates presumptive evidence of a compound







Page 1 of 1

Client Sample ID: MW-7 Lab Sample ID: JC1603

Matrix:

JC16038-2

AQ - Ground Water SW846 8270D BY SIM SW846 3510C Date Sampled: 03/09/16 Date Received: 03/11/16

Q

Percent Solids: n/a

Method: Project:

BMSMC, Building 5 Area, PR

 File ID
 DF
 Analyzed
 By
 Prep Date
 Prep Batch
 Analytical Batch

 Run #1
 4P15637.D
 1
 03/17/16
 JJ
 03/15/16
 OP92078A
 E4P804

Run #2

Initial Volume Final Volume
Run #1 950 ml 1.0 ml

Run #2

CAS No. RL MDL Compound Result Units 91-20-3 Naphthalene ND 0.014 0.11 ug/l 123-91-1 1,4-Dioxane 0.056 1.72 0.11 ug/l CAS No. Surrogate Recoveries Run#1 Run# 2 Limits

 4165-60-0
 Nitrobenzene-d5
 79%
 24-125%

 321-60-8
 2-Fluorobiphenyl
 74%
 19-127%

 1718-51-0
 Terphenyl-d14
 89%
 10-119%



ND = Not detected

MDL = Method Detection Limit

J = Indicates an estimated value

RL = Reporting Limit

B = Indicates analyte found in associated method blank N = Indicates presumptive evidence of a compound

E = Indicates value exceeds calibration range



SGS Accutest

Report of Analysis

Page 1 of 1

Client Sample ID: MW-7

Lab Sample ID: JC16038-2

Matrix; Method: AQ - Ground Water

SW846-8015C (DAI)

Date Sampled: 03/09/16 Date Received:

03/11/16

Percent Solids: n/a

Project: BMSMC, Building 5 Area, PR

Analytical Batch File ID DF Analyzed By Prop Date Prep Batch Run #1 GH103822.D 03/18/16 XPL **GGH5213** 1 n/a n/a Run #2

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units	Q
64-17-5	Ethanol	ND	100	55	ug/l	
78-83-1	Isobutył Alcohol	ND	100	36	ug/l	
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l	
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l	
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l	
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l	
67-56-1	Methanol	ND	200	71	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
111-27-3	Hexanol	93%		56-1	45%	
111-27-3	Hexanol	94%		56-1	45%	





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



SGS Accutest

Report of Analysis

Page 1 of 2

Client Sample ID: MW-3

Lab Sample ID: JC16038-3

Matrix: Method:

Project:

AQ - Ground Water SW846 8260C

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Q

J

J

J

J

Percent Solids: n/a

	File ID	DF	Analyzed	Ву	Prep Date	Prep Batch	Analytical Batch
Run #1	U204147.D	1	03/15/16	NH	n/a	n/a	VU9383
D., a #2							

Run #2

Purge Volume

Run #1 5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	0.32	0.50	0.24	ug/I
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	0.36	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	4.0	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	0.27	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	ND	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	ND	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34 - 3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	0.45	1.0	0.27	ug/l
76-13-1	Freon 113	ND	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l

fael Infante

ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 2

Client Sample ID: MW-3

Lab Sample ID: JC16038-3

Matrix: Method: AQ - Ground Water SW846 8260C

Project:

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16

Date Received: 03/11/16

Percent Solids: n/a

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	18.6	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	4.9	5.0	0.22	ug/l	J
1634-04-4	Methyl Tert Butyl Ether	ND	1.0	0.24	ug/l	-
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1:0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	0.42	1.0	0.16	ug/l	J
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	_
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	0.61	1.0	0.38	ug/l	j
95-47-6	o-Xylene	0.98	1.0	0.17	ug/l	Ī
1330-20-7	Xylene (total)	1.6	1.0	0.17	ug/l	_
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limi	ts	
1868-53-7	Dibromofluoromethane	105%		76-12	20%	
17060-07-0	1,2-Dichloroethane-D4	108%		73-12	2%	

99%

104%





2037-26-5

460-00-4

Toluene-D8

4-Bromofluorobenzene

84-119%

78-117%



MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Client Sample ID: Lab Sample ID:

MW-3 JC16038-3

Matrix:

AQ - Ground Water

Method: Project:

SW846 8270D SW846 3510C

BMSMC, Building 5 Area, PR

Date Sampled: Date Received:

Q

J

03/09/16 03/11/16

Percent Solids: n/a

		_					
71 87	File ID	DF	Analyzed	Ву	Prep Date	Prep Batch	Analytical Batch
Run #1	F155579.D	1	03/16/16	SD	03/15/16	OP92078	EF6543

Run #2

Initial Volume 1000 ml

Final Volume 1.0 ml

Run #1 Run #2

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Unit
95-57-8	2-Chlorophenol	ND	5.0	0.93	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.0	1.4	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.0	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.0	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	10	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.0	0.87	ug/l
95-48-7	2-Methylphenol	ND	2.0	0.82	ug/l
	3&4-Methylphenoi	ND	2.0	0.67	ug/l
88-75-5	2-Nitrophenol	ND	5.0	1.4	ug/l
100-02-7	4-Nitrophenol	ND	10	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.0	1.4	ug/l
108-95-2	Phenol	ND	2.0	0.31	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.0	1.4	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.0	1.5	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.0	1.4	ug/l
83-32-9	Acenaphthene	0.67	1.0	0.29	ug/l
208-96-8	Acenaphthylene	ND	1.0	0.24	ug/l
98-86-2	Acetophenone	ND	2.0	0.28	ug/l
120-12-7	Anthracene	ND	1.0	0.25	ug/l
1912-24-9	Atrazine	ND	2.0	0.42	ug/l
100-52-7	Benzaldehyde	ND	5.0	0.34	ug/l
56-55-3	Benzo(a)anthracene	ND	1.0	0.32	ug/l
50-32-8	Benzo(a)pyrene	ND	1.0	0.33	ug/l
205-99-2	Benzo(b) fluoranthene	ND	1.0	0.32	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.0	0.41	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.0	0.37	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.0	0.37	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.0	0.27	ug/l
92-52-4	1,1'-Biphenyl	ND	1.0	0.26	ug/l
91-58-7	2-Chloronaphthalene	ND	2.0	0.30	ug/l
106-47-8	4-Chloroaniline	ND	5.0	0.23	ug/l
86-74-8	Carbazole	ND	1.0	0.29	ug/l



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 3

Client Sample ID: MW-3

Lab Sample ID: JC16038-3 Matrix:

AQ - Ground Water Method: SW846 8270D SW846 3510C Project:

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

Q

ABN TCL List (SOM0 1.1)

CAS No.	CAS No. Compound		RL	MDL	Unit
105-60-2	Caprolactam	ND	2.0	0.43	ug/l
218-01-9	Chrysene	ND	1.0	0.35	ug/l
111-91-1	bis(2-Chloroethoxy)methane	ND	2.0	0.26	ug/l
111-44-4	bis(2-Chloroethyl)ether	ND	2.0	0.34	ug/l
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.0	0.28	ug/l
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.0	0.27	ug/l
121-14-2	2,4-Dinitrotoluene	ND	1.0	0.26	ug/l
606-20-2	2,6-Dinitrotoluene	ND	1.0	0.32	ug/I
91-94-1	3,3'-Dichlorobenzidine	ND	2.0	0.53	ug/l
53-70-3	Dibenzo(a,h)anthracene	ND	1.0	0.37	ug/l
132-64-9	Dibenzofuran	ND	5.0	0.27	ug/l
84-74-2	Di-n-butyl phthalate	ND	2.0	0.79	ug/l
117-84-0	Di-n-octyl phthalate	ND	2.0	0.29	ug/l
84-66-2	Diethyl phthalate	ND	2.0	0.24	ug/l
131-11-3	Dimethyl phthalate	ND	2.0	0.31	ug/l
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.0	0.77	ug/l
206-44-0	Fluoranthene	ND	1.0	0.23	ug/l
86-73-7	Fluorene	1.2	1.0	0.29	ug/l
118-74-1	Hexachlorobenzene	ND	1.0	0.42	ug/l
87-68-3	Hexachlorobutadiene	ND	1.0	0.36	ug/l
77-47-4	Hexachlorocyclopentadiene	ND	10	0.29	ug/l
67-72-1	Hexachloroethane	ND	2.0	0.22	ug/l
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.0	0.38	ug/l
78-59-1	Isophorone	ND	2.0	0.29	ug/I
91-57-6	2-Methylnaphthalene	52.4	1.0	0.29	ug/l
88-74-4	2-Nitroaniline	ND	5.0	0.21	ug/l
99-09-2	3-Nitroaniline	ND	5.0	0.24	ug/l
100-01-6	4-Nitroaniline	ND	5.0	0.34	ug/l
91-20-3	Naphthalene	3.6	1.0	0.28	ug/l
98-95-3	Nitrobenzene	ND	2.0	0.46	ug/l
621-64-7	N-Nitroso-di-n-propylamine	ND	2.0	0.31	ug/l
86-30-6	N-Nitrosodiphenylamine	ND	5.0	0.29	ug/l
85-01-8	Phenanthrene	ND	1.0	0.23	ug/l
129-00-0	Pyrene	ND	1.0	0.34	ug/l
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.0	0.36	ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its
367-12-4	2-Fluorophenol	42%		14-8	8%
4405 00 0	701 1 17				



ND = Not detected

4165-62-2

MDL = Method Detection Limit

32%

RL = Reporting Limit

E = Indicates value exceeds calibration range

Phenol-d5

J = Indicates an estimated value

10-110%

B = Indicates analyte found in associated method blank



Project:

Report of Analysis

Page 3 of 3

Client Sample ID: MW-3

Lab Sample ID: JC16038-3

Matrix: AQ - Ground Water Method: SW846 8270D SW

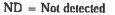
SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limits
118-79-6	2,4,6-Tribromophenol	71%		39-149%
4165-60-0	Nitrobenzene-d5	67%		32-128%
321-60-8	2-Fluorobiphenyl	64%		35-119%
1718-51-0	Terphenyl-d14	68%		10-126%





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 1 of 1

Client Sample ID: MW-3 Lab Sample ID: JC16038-3

Initial Volume

Terphenyl-d14

Matrix:

AQ - Ground Water

Method: Project:

1718-51-0

SW846 8270D BY SIM SW846 3510C

BMSMC, Building 5 Area, PR

Final Volume

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

	File ID	DF	Analyzed	Ву	Prep Date	Prep Batch	Analytical Batch
Run #1	4P15638.D	1	03/17/16	IJ	03/15/16	OP92078A	E4P804
Run #2							

Run #1 Run #2	1000 ml 1.0	0 ml				
CAS No.	Compound	Result	RL	MDL	Units	Q
91-20-3	Naphthalene	3.57	0.10	0.013	ug/l	
123-91-1	1,4-Dioxane	ND	0.10	0.053	ug/l	
CAS No.	Surrogate Recover	ries Run#1	Run# 2	2 Lim	its	
4165-60-0	Nitrobenzene-d5	63%		24-1	25%	
321-60-8	2-Fluorobiphenyl	68%		19-1	27%	

75%



10-119%



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank N = Indicates presumptive evidence of a compound

Page 1 of 1

Client Sample ID: MW-3

Lab Sample ID: JC16038-3

Matrix: Method:

Project:

AQ - Ground Water

SW846-8015C (DAI) BMSMC, Building 5 Area, PR Date Sampled: 03/09/16 Date Received:

03/11/16

Percent Solids: n/a

File ID DF Analyzed Prep Date Prep Batch **Analytical Batch** By GH103823.D Run #1 1 03/18/16 XPL **GGH5213** n/a n/a Run #2

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units	Q
64-17-5	Ethanol	ND	100	55	ug/l	
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l	
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l	
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l	
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l	
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l	
67-56-1	Methanol	ND	200	71	ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its	
111-27-3	Hexanol	93%		56-1	45%	
111-27-3	Hexanol	91%		56-1	45%	





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



SGS Accutest

Report of Analysis

Page 1 of 2

Client Sample ID: MW-5

Lab Sample ID: JC16038-4 Matrix: AQ - Ground Water

Method: SW846 8260C Project:

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

	File ID	DF	Analyzed	Ву	Prep Date	Prep Batch	Analytical Batch
Run #1	U204148.D	1	03/15/16	NH	n/a	n/a	VU9383
Run #2	A221006.D	20	03/17/16	NH	n/a	n/a	VA8370

Purge Volume Run #1 $5.0 \, ml$ 5.0 ml Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
67-64-1	Acetone	ND	10	3.3	ug/l	
71-43-2	Benzene	7.4	0.50	0.24	ug/l	
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l	
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l	
75-25-2	Bromoform	ND	1.0	0.23	ug/l	
74-83-9	Bromomethane	ND	2.0	0.42	ug/l	
78-93- 3	2-Butanone (MEK)	ND	10	5.6	ug/l	
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l	
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l	
108-90-7	Chlorobenzene	0.36	1.0	0.19	ug/l	J
75-00-3	Chloroethane	ND	1.0	0.34	ug/l	
67-66-3	Chloroform	ND	1.0	0.19	ug/l	
74-87-3	Chloromethane	ND	1.0	0.41	ug/l	
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l	
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/I	
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l	
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l	
95-50-1	1,2-Dichlorobenzene	0.48	1.0	0.19	ug/l	J
541-73-1	1,3-Dichlorobenzene	ND	1.0	0.23	ug/l	
106-46-7	1,4-Dichlorobenzene	ND	1.0	0.27	ug/l	
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l	
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l	
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l	
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l	
156-59-2	cis-1,2-Dichloroethene	0.29	1.0	0.27	ug/l	J
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l	_
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l	
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l	
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l	
100-41-4	Ethylbenzene	1040 ª	20	5.4	ug/l	
76-13-1	Freon 113	ND	5.0	0.52	ug/l	
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l	



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 2

Client Sample ID: Lab Sample ID:

MW-5JC16038-4

Matrix: Method: AQ - Ground Water

Project:

SW846 8260C

BMSMC, Building 5 Area, PR

Date Sampled: Date Received:

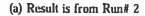
03/09/16

03/11/16

Percent Solids:

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	20.8	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	11.8	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	0.25	1.0	0.16	ug/l	J
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	_
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	2480 a	20	7.5	ug/l	
95-47-6	o-Xylene	0.42	1.0	0.17	ug/l	J
1330-20-7	Xylene (total)	2480 a	20	3.3	ug/l	_
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
1868-53-7	Dibromofluoromethane	105%	104%	76-1	20%	
17060-07-0	1,2-Dichloroethane-D4	107%	102%	73-1	22%	
2037-26-5	Toluene-D8	100%	100%	84-1	19%	11
460-00-4	4-Bromofluorobenzene	100%	100%	78-1	17%	99







MDL = Method Detection Limit

RL = Reporting Limit



E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Client Sample ID: MW-5 Lab Sample ID:

JC16038-4

Matrix: Method: AQ - Ground Water

Date Sampled: Date Received:

03/09/16 03/11/16

Project:

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Percent Solids: n/a

Q

	File ID	DF	Analyzed	Ву	Prep Date	Prep Batch	Analytical Batch
Run #1	F155580.D	1	03/16/16	SD	03/15/16	OP92078	EF6543
T #0							

Run #2

Initial Volume Final Volume 950 ml

Run #1 Run #2 1.0 ml

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Unit
95-57-8	2-Chlorophenol	ND	5.3	0.98	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.3	1.5	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.1	1.3	ug/l
105-67-9	2,4-Dimethylphenol	5.7	5.3	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	11	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.3	0.92	ug/l
95-48-7	2-Methylphenol	ND	2.1	0.86	ug/l
	3&4-Methylphenol	ND	2.1	0.71	ug/l
88-75-5	2-Nitrophenol	ND	5.3	1.5	ug/l
100-02-7	4-Nitrophenol	ND	11	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.3	1.5	ug/l
108-95-2	Phenol	ND	2.1	0.33	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.3	1.5	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.3	1.6	ug/l
88-06-2	2,4,6-Trichtorophenol	ND	5.3	1.5	ug/l
83-32-9	Acenaphthene	ND	1.1	0.30	ug/l
208-96-8	Acenaphthylene	ND	1.1	0.25	ug/l
98-86-2	Acetophenone	ND	2.1	0.29	ug/l
120-12-7	Anthracene	ND	1.1	0.26	ug/l
1912-24-9	Atrazine	ND	2.1	0.44	ug/l
100-52-7	Benzaldehyde	ND	5.3	0.35	ug/l
56-55-3	Benzo(a)anthracene	ND	1.1	0.33	ug/l
50-32-8	Benzo(a) pyrene	ND	1.1	0.35	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.1	0.33	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.1	0.43	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.1	0.39	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.1	0.39	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.1	0.29	ug/l
92-52-4	1,1'-Biphenyl	ND	1.1	0.27	ug/l
91-58-7	2-Chloronaphthalene	ND	2.1	0.31	ug/l
106-47-8	4-Chloroaniline	ND	5.3	0.24	ug/l
86-74-8	Carbazole	ND	1.1	0.31	ug/l

tael Infante Méndez-

ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 3

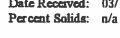
Client Sample ID: MW-5 Lab Sample ID: JC16038-4

Matrix: AQ - Ground Water

Method: SW846 8270D SW846 3510C Project: BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received:

03/11/16



ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Units	Q
105-60-2	Caprolactam	ND	2.1	0.45	ug/l	
218-01-9	Chrysene	ND	1.1	0.36	ug/l	
111-91-1	bis(2-Chloroethoxy)methane	ND	2.1	0.27	ug/l	
111-44-4	bis(2-Chloroethyl)ether	ND	2.1	0.36	ug/l	
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.1	0.30	ug/l	
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.1	0.29	ug/l	
121-14-2	2,4-Dinitrotoluene	ND	1.1	0.28	ug/l	
606-20-2	2,6-Dinitrotoluene	ND	1.1	0.34	ug/l	
91-94-1	3,3'-Dichlorobenzidine	ND	2.1	0.56	ug/l	
53-70-3	Dibenzo (a,h) anthracene	ND	1.1	0.38	ug/l	
132-64-9	Dibenzofuran	ND	5.3	0.29	ug/l	
84-74-2	Di-n-butyl phthalate	ND	2.1	0.83	ug/l	
117-84-0	Di-n-octyl phthalate	ND	2.1	0.30	ug/l	
84-66-2	Diethyl phthalate	ND	2.1	0.26	ug/l	
131-11-3	Dimethyl phthalate	ND	2.1	0.33	ug/l	
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.1	0.81	ug/l	
206-44-0	Fluoranthene	ND	1.1	0.24	ug/l	
86-73-7	Fluorene	ND	1.1	0.31	ug/l	
118-74-1	Hexachlorobenzene	ND	1.1	0.44	ug/l	
87-68-3	Hexachlorobutadiene	ND	1.1	0.38	ug/l	
77-47-4	Hexachlorocyclopentadiene	ND	11	0.31	ug/l	
67-72-1	Hexachloroethane	ND	2.1	0.23	ug/l	
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.1	0.40	ug/l	
78-59-1	Isophorone	ND	2.1	0.30	ug/l	
91-57-6	2-Methylnaphthalene	0.63	1.1	0.30	ug/l	J
88-74-4	2-Nitroaniline	ND	5.3	0.22	ug/l	-
99-09-2	3-Nitroaniline	ND	5.3	0.25	ug/l	
100-01-6	4-Nitroaniline	ND	5.3	0.36	ug/l	
91-20-3	Naphthalene	2.7	1.1	0.30	ug/l	
98-95-3	Nitrobenzene	ND	2.1	0.49	ug/l	1222
621-64-7	N-Nitroso-di-n-propylamine	ND	2.1	0.33	ug/l	LIDAD
86-30-6	N-Nitrosodiphenylamine	ND	5.3	0.31	ug/l	SAE ISOCIA
85-01-8	Phenanthrene	ND	1.1	0.24	ug/l	1 3
129-00-0	Pyrene	ND	1.1	0.35	ug/l	[5] I fael I
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.1	0.38	ug/l	Ména Ména
						C 1 1 2
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	14
						EMICO LIC
367-12-4	2-Fluorophenol	42%		14-8		OLIU
4165-62-2	Phenol-d5	32%		10-1	10%	

ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 3 of 3

Client Sample ID: MW-5

JC16038-4 Lab Sample ID:

Matrix: Method:

Project:

AQ - Ground Water

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a



ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limits
118-79-6	2,4,6-Tribromophenol	83%		39-149%
4165-60-0	Nitrobenzene-d5	71%		32-128%
321-60-8	2-Fluorobiphenyl	70%		35-119%
1718-51-0	Terphenyl-d14	73%		10-126%





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 1 of 1

Client Sample ID: Lab Sample ID:

MW-5 JC16038-4

Matrix: Method: AQ - Ground Water

SW846 8270D BY SIM SW846 3510C

Date Sampled: Date Received:

03/09/16 03/11/16

Percent Solids: n/a

Project:

BMSMC, Building 5 Area, PR

File ID DF Analyzed Ву Prep Date Prep Batch **Analytical Batch** Run #1 4P15639.D 1 03/17/16 IJ 03/15/16 OP92078A E4P804 Run #2

	Initial Volume	Final Volume
Run #1	950 ml	1.0 ml

Run #2

CAS No.	Compound	Result	RL	MDL	Units	Q
91-20-3 123-91-1	Naphthalene 1,4-Dioxane	2.48 0.734	0.11 0.11	0.014 0.056	ug/l ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limi	ts	
4165-60-0 321-60-8 1718-51-0	Nitrobenzene-d5 2-Fluorobiphenyl Terphenyl-d14	73% 73% 80%		24-12 19-12 10-11	27%	



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

Page 1 of 1

Client Sample ID: MW-5 Lab Sample ID:

Matrix: Method: JC16038-4 AQ - Ground Water

Project:

SW846-8015C (DAI)

BMSMC, Building 5 Area, PR

Date Sampled: Date Received: 03/11/16

03/09/16

Percent Solids: n/a

Run #1	File ID	DF	Analyzed	By	Prep Date	Prep Batch	Analytical Batch
Run #2	GH103824.D	1	03/18/16	XPL	n/a	n/a	GGH5213
Run #2							

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units	Q
64-17-5	Ethanol	ND	100	55	ug/l	
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l	
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l	
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l	
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l	
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l	
67-56-1	Methanol	ND	200	71	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
111-27-3	Hexanol	100%		56-1	45%	
111-27-3	Hexanol	97%		56-1	45%	1





MDL = Method Detection Limit

RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



E = Indicates value exceeds calibration range

SGS Accutest

Report of Analysis

Page 1 of 2

Client Sample ID: MW-16

Lab Sample ID: JC16038-5

Matrix: Method: AQ - Ground Water SW846 8260C

Date Sampled: Date Received: 03/11/16

Q

J

J

03/09/16

Percent Solids: n/a

Project:

BMSMC, Building 5 Area, PR

File ID DF Analyzed By Prep Date Prep Batch **Analytical Batch** Run #1 A221005.D NH 1 03/17/16 VA8370 n/a n/a

Run #2

Purge Volume

Run #1 5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Unite
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	0.34	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	12.2	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	0.49	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	2.5	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	ND	1.0	0.27	ug/l
76-13-1	Freon 113	58.5	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l
					_



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank





Page 2 of 2

Client Sample ID: Lab Sample ID:

MW-16 JC16038-5

Matrix:

AQ - Ground Water SW846 8260C

Method: Project:

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a



VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	ND	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	ND	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	NĐ	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	ND	1.0	0.38	ug/l	
95-47-6	o-Xylene	ND	1.0	0.17	ug/l	
1330-20-7	Xylene (total)	ND	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limi	ts	
1868-53-7	Dibromofluoromethane	103%		76-17	20%	
17060-07-0	1,2-Dichloroethane-D4	100%		73-17	22%	
2037-26-5	Toluene-D8	100%		84-11	19%	
460-00-4	4-Bromofluorobenzene	99%		78-1	17%	/



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Client Sample ID: MW-16

Lab Sample ID: JC16038-5

Matrix: AQ - Ground Water Method: SW846 8270D SW846 3510C Project:

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

Q

File ID DF Analyzed **Analytical Batch** By Prep Date Prep Batch Run #1 F155581.D 1 03/16/16 SĎ 03/15/16 OP92078 EF6543

Run #2

Initial Volume Final Volume

Run #1 950 ml 1.0 ml

Run #2

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Unite
95-57-8	2-Chlorophenol	ND	5.3	0.98	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.3	1.5	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.1	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.3	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	11	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.3	0.92	ug/l
95-48-7	2-Methylphenol	ND	2.1	0.86	ug/l
	3&4-Methylphenol	ND	2.1	0.71	ug/l
88-75-5	2-Nitrophenol	ND	5.3	1.5	ug/l
100-02-7	4-Nitrophenol	ND	11	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.3	1.5	ug/l
108-95-2	Phenol	ND	2.1	0.33	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.3	1.5	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.3	1.6	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.3	1.5	ug/l
83-32-9	Acenaphthene	ND	1.1	0.30	ug/l
208-96-8	Acenaphthylene	ND	1.1	0.25	ug/l
98-86-2	Acetophenone	ND	2.1	0.29	ug/l
120-12-7	Anthracene	ND	1.1	0.26	ug/l
1912-24-9	Atrazine	ND	2.1	0.44	ug/l
100-52-7	Benzaldehyde	ND	5.3	0.35	ug/l
56-55-3	Benzo(a)anthracene	ND	1:1	0.33	ug/l
50-32-8	Benzo(a)pyrene	ND	1.1	0.35	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.1	0.33	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.1	0.43	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.1	0.39	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.1	0.39	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.1	0.29	ug/l
92-52-4	1,1'-Biphenyl	ND	1.1	0.27	ug/l
91-58-7	2-Chloronaphthalene	ND	2.1	0.31	ug/l
106-47-8	4-Chloroaniline	ND	5.3	0.24	ug/l
86-74-8	Carbazole	ND	1.1	0.31	ug/l

fael Infante Méndez

ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 3

Client Sample ID: Lab Sample ID:

MW-16 JC16038-5

Matrix:

AQ - Ground Water

Method: Project:

SW846 8270D SW846 3510C

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

Q

ABN TCL List (SOM0 1.1)

	,				
CAS No.	Compound	Result	RL	MDL	Units
105-60-2	Caprolactam	ND	2.1	0.45	ug/l
218-01-9	Chrysene	ND	1.1	0.36	ug/l
111-91-1	bis(2-Chloroethoxy)methane	ND	2.1	0.27	ug/l
111-44-4	bis(2-Chloroethyl)ether	ND	2.1	0.36	ug/l
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.1	0.30	ug/l
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.1	0.29	ug/l
121-14-2	2,4-Dinitrotoluene	ND	1.1	0.28	ug/l
606-20-2	2,6-Dinitrotoluene	ND	1.1	0.34	ug/l
91-94-1	3,3'-Dichlorobenzidine	ND	2.1	0.56	ug/l
53-70-3	Dibenzo(a,h)anthracene	ND	1.1	0.38	ug/l
132-64-9	Dibenzofuran	ND	5.3	0.29	ug/l
84-74-2	Di-n-butyl phthalate	ND	2.1	0.83	ug/l
117-84-0	Di-n-octyl phthalate	ND	2.1	0.30	ug/l
84-66-2	Diethyl phthalate	ND	2.1	0.26	ug/l
131-11-3	Dimethyl phthalate	ND	2.1	0.33	ug/l
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.1	0.81	ug/l
206-44-0	Fluoranthene	ND	1.1	0.24	ug/l
86-73-7	Fluorene	ND	1.1	0.31	ug/l
118-74-1	Hexachlorobenzene	ND	1.1	0.44	ug/l
87-68-3	Hexachlorobutadiene	ND	1.1	0.38	ug/l
77-47-4	Hexachlorocyclopentadiene	ND	11	0.31	ug/l
67-72-1	Hexachloroethane	ND	2.1	0.23	ug/l
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.1	0.40	ug/l
78-59 -1	Isophorone	ND	2.1	0.30	ug/l
91-57-6	2-Methylnaphthalene	ND	1.1	0.30	ug/l
88-74-4	2-Nitroaniline	ND	5.3	0.22	ug/l
99-09-2	3-Nitroaniline	ND	5.3	0.25	ug/l
100-01-6	4-Nitroaniline	ND	5.3	0.36	ug/l
91-20-3	Naphthalene	ND	1.1	0.30	ug/l
98-95-3	Nitrobenzene	ND	2.1	0.49	ug/l
621-64-7	N-Nitroso-di-n-propylamine	ND	2.1	0.33	ug/l
86-30-6	N-Nitrosodiphenylamine	ND	5.3	0.31	ug/l
85-01-8	Phenanthrene	ND	1.1	0.24	ug/l
129-00-0	Ругепе	ND	1.1	0.35	ug/l
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.1	0.38	ug/l
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limits	
367-12-4	2-Fluorophenol	42%		14-88%	
4165-62-2	Phenol-d5	32%		10-110%	



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

E = Indicates value exceeds calibration range

SGS Accutest

Report of Analysis

Page 3 of 3

Client Sample ID: MW-16 Lab Sample ID: JC16038-5 Matrix:

AQ - Ground Water

Date Sampled: Date Received: 03/11/16

03/09/16

Method: Project:

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Percent Solids: n/a

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limits
118-79-6	2,4,6-Tribromophenol	77%		39-149%
4165-60-0	Nitrobenzene-d5	72%		32-128%
321-60-8	2-Fluorobiphenyl	66%		35-119%
1718-51-0	Terphenyl-d14	71%		10-126%





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



By

IJ

Client Sample ID: Lab Sample ID:

MW-16 JC16038-5

Matrix:

Method: Project:

AQ - Ground Water

DF

SW846 8270D BY SIM SW846 3510C

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16

Date Received: 03/11/16

Percent Solids: n/a

Prep Batch

OP92078A

Analytical Batch E4P804

Run #1 Run #2

Run #1

Run #2

91-20-3

123-91-1

CAS No.

Final Volume Initial Volume

1.0 ml

950 ml

CAS No. Compound

File ID

4P15640.D

Naphthalene 1,4-Dioxane

Surrogate Recoveries

4165-60-0 Nitrobenzene-d5 321-60-8 2-Fluorobiphenyl 1718-51-0 Terphenyl-d14

Result

ND

0.388

Run#1

74%

73%

79%

Analyzed

03/17/16

0.11 0.11

Run# 2

RL

0.014 0.056

MDL

ug/l ug/l

Limits

Units

Q

Prep Date

03/15/16

24-125% 19-127%

10-119%





ND = Not detected

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

SGS Accutest

Report of Analysis

Page 1 of 1

Client Sample ID: MW-16 Lab Sample ID: JC16038-5

Matrix: Method:

Project:

AQ - Ground Water

SW846-8015C (DAI) BMSMC, Building 5 Area, PR Date Sampled: Date Received:

03/09/16 03/11/16

Percent Solids: n/a

File ID DF **Analytical Batch** Analyzed By Prep Date Prep Batch Run #1 XPL GH103825.D 03/18/16 GGH5213 n/a n/a Run #2

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units	Q
64-17-5	Ethanol	ND	100	55	ug/l	
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l	
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l	
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l	
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l	
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l	
67-56-1	Methanol	ND	200	71	ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limits		
111-27-3	Hexanol	96%		56-1	45%	
111-27-3	Hexanol	93%		56-1	45%	





MDL = Method Detection Limit

RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



E = Indicates value exceeds calibration range

Report of Analysis

By

NH

Prep Date

n/a

Page 1 of 2

Client Sample ID:

MW-16D

Lab Sample ID:

File ID

U204161.D

JC16038-6 AQ - Ground Water Date Sampled: 03/09/16

Matrix: Method:

SW846 8260C

Date Received: 03/11/16

Project:

DF

Percent Solids: n/a

n/a

Q

J

J

BMSMC, Building 5 Area, PR

Analyzed

03/15/16

Prep Batch

Analytical Batch VU9383

Run #1 Run #2

Purge Volume

Run #1

Run #2

 $5.0 \, ml$

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Unit
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	0.32	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	12.7	1,0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	0.46	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	2.5	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	ND	1.0	0.27	ug/l
76-13-1	Freon 113	44.0	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l
					_



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 2

Client Sample ID: Lab Sample ID:

MW-16D JC16038-6

Matrix:

AQ - Ground Water

Method:

SW846 8260C

Project:

BMSMC, Building 5 Area, PR

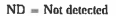
Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	ND	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	ND	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	ND	1.0	0.38	ug/l	
95-47-6	o-Xylene	ND	1.0	0.17	ug/l	
1330-20-7	Xylene (total)	ND	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its	
1868-53-7	Dibromofluoromethane	106%		76-1	20%	
17060-07-0	1,2-Dichloroethane-D4	107%		73-1	22%	
2037-26-5	Toluene-D8	99%		84-1	19%	
460-00-4	4-Bromofluorobenzene	103%		78-1	17%	0





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Client Sample ID: Lab Sample ID:

MW-16D JC16038-6

AQ - Ground Water

Matrix: Method:

SW846 8270D SW846 3510C

Project:

BMSMC, Building 5 Area, PR

Date Sampled:

Q

J

03/09/16

Date Received:

03/11/16

Percent Solids:

n/a

î -							
	File ID	DF	Analyzed	By	Prep Date	Prep Batch	Analytical Batch
Run #1	F155582.D	1	03/16/16	SD	03/15/16	OP92078	EF6543

Run #2

Initial Volume Final Volume

Run #1

970 ml

1.0 ml

Run #2

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Unit
95-57-8	2-Chlorophenol	ND	5.2	0.96	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.2	1.4	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.1	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.2	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	10	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.2	0.90	ug/l
95-48-7	2-Methylphenol	ND	2.1	0.84	ug/l
	3&4-Methylphenol	ND	2.1	0.69	ug/l
88-75-5	2-Nitrophenol	ND	5.2	1.5	ug/l
100-02-7	4-Nitrophenol	ND	10	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.2	1.5	ug/l
108-95-2	Phenol	ND	2.1	0.32	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.2	1.5	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.2	1.5	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.2	1.5	ug/l
83-32-9	Acenaphthene	0.44	1.0	0.29	ug/l
208-96-8	Acenaphthylene	ND	1.0	0.25	ug/l
98-86-2	Acetophenone	ND	2.1	0.28	ug/l
120-12-7	Anthracene	ND	1.0	0.25	ug/l
1912-24-9	Atrazine	ND	2.1	0.43	ug/l
100-52-7	Benzaldehyde	ND	5.2	0.35	ug/l
56-55-3	Benzo(a)anthracene	ND	1.0	0.32	ug/l
50-32-8	Вепхо(а)ругепе	ND	1.0	0.34	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.0	0.33	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.0	0.42	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.0	0.38	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.1	0.38	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.1	0.28	ug/l
92-52-4	1.1 -Biphenyl	ND	1.0	0.26	ug/l
91-58-7	2-Chloronaphthalene	ND	2.1	0.31	ug/l
106-47-8	4-Chloroaniline	ND	5.2	0.24	ug/l
86-74-8	Carbazole	ND	1.0	0.30	ug/l



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 3

Client Sample ID:

MW-16D JC16038-6

Lab Sample ID:

AQ - Ground Water

Matrix: Method:

Project:

SW846 8270D SW846 3510C

B

BMSMC, Building 5 Area, PR

Date Sampled: Date Received:

03/09/16 03/11/16

Percent Solids: n/a



ABN TCL List (SOM0 1.1)

111111101	Em (BOING 1.1)						
CAS No.	Compound	Result	RL	MDL	Units	Q	
105-60-2	Caprolactam	ND	2.1	0.44	ug/l		
218-01-9	Chrysene	ND	1.0	0.36	ug/l		
111-91-1	bis(2-Chloroethoxy)methane	ND	2.1	0.27	ug/l		
111-44-4	bis(2-Chloroethyl)ether	ND	2.1	0.35	ug/l		
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.1	0.29	ug/l		
7005-72-3		ND	2.1	0.28	ug/l		
121-14-2	2,4-Dinitrotoluene	ND	1.0	0.27	ug/l		
606-20-2	2,6-Dinitrotoluene	ND	1.0	0.33	ug/l		
91-94-1	3,3'-Dichlorobenzidine	ND	2.1	0.55	ug/l		
53-70-3	Dibenzo (a,h) anthracene	ND	1.0	0.38	ug/l		
132-64-9	Dibenzofuran	ND	5.2	0.28	ug/l		
84-74-2	Di-n-butyl phthalate	ND	2.1	0.81	ug/l		
117-84-0	Di-n-octyl phthalate	ND	2.1	0.30	ug/l		
84-66-2	Diethyl phthalate	ND	2.1	0.25	ug/l		
131-11-3	Dimethyl phthalate	ND	2.1	0.32	ug/l		
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.1	0.79	ug/l		
206-44-0	Fluoranthene	ND	1.0	0.24	ug/l		
86-73-7	Fluorene	0.49	1.0	0.30	ug/l	J	
118-74-1	Hexachlorobenzene	ND	1.0	0.44	ug/l		
87-68-3	Hexachlorobutadiene	ND	1.0	0.38	ug/l		
77-47-4	Hexachlorocyclopentadiene	ND	10	0.30	ug/l		
67-72-1	Hexachloroethane	ND	2.1	0.23	ug/l		
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.0	0.40	ug/l		
78-59-1	Isophorone	ND	2.1	0.29	ug/l		
91-57-6	2-Methylnaphthalene	3.4	1.0	0.30	ug/l		
88-74-4	2-Nitroaniline	ND	5.2	0.22	ug/l		
99-09-2	3-Nitroaniline	ND	5.2	0.25	ug/l		
100-01-6	4-Nitroaniline	ND	5.2	0.35	ug/l		
91-20-3	Naphthalene	2.3	1.0	0.29	ug/l		
98-95-3	Nitrobenzene	ND	2.1	0.48	ug/l		2
621-64-7	N-Nitroso-di-n-propylamine	ND	2.1	0.32	ug/l		
86-30-6	N-Nitrosodiphenylamine	ND	5.2	0.30	ug/l		123
85-01-8	Phenanthrene	0.61	1.0	0.24	ug/l	J	1-5
129-00-0	Pyrene	ND	1.0	0.35	ug/l		E2714
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.1	0.37	ug/l		1.2
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its		
367-12-4	2-Fluorophenol	55%		14-8	8%		
4105 00 0	Diagonal ar	2007		10.5	100/		

ND = Not detected

4165-62-2

MDL = Method Detection Limit

39%

RL = Reporting Limit

E = Indicates value exceeds calibration range

Phenol-d5

J = Indicates an estimated value

10-110%

B = Indicates analyte found in associated method blank

Méndez



Report of Analysis

Page 3 of 3

Client Sample ID: MW-16D Lab Sample ID: JC16038-6 Matrix:

AQ - Ground Water

Date Sampled: 03/09/16 Date Received: 03/11/16 Percent Solids:

Method: Project:

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limits
118-79-6	2,4,6-Tribromophenol	97%		39-149%
4165-60-0	Nitrobenzene-d5	97%		32-128%
321-60-8	2-Fluorobiphenyl	89%		35-119%
1718-51-0	Terphenyl-d14	86%		10-126%





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



By

IJ

Prep Date

03/15/16

Client Sample ID: Lab Sample ID:

MW-16D JC16038-6

Matrix: Method:

Project:

File ID

970 ml

4P15641.D

AQ - Ground Water SW846 8270D BY SIM SW846 3510C

DF

1

BMSMC, Building 5 Area, PR

Analyzed

03/17/16

Date Sampled:

03/09/16 03/11/16

Date Received: Percent Solids:

n/a

OP92078A

Q

Prep Batch **Analytical Batch**

E4P804

Run #1 Run #2

> Initial Volume Final Volume

Run #1 Run #2 1.0 ml

CAS No.	Compound	Result	RL	MDL	Units
91-20-3	Naphthalene	2.31	0.10	0.014	ug/l
123-91-1	1,4-Dioxane	0.925	0.10	0.055	ug/l

123-91-1	I,4-Dioxane	0.925	0.10	0.055 ug	į/Ι
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limits	
4165-60-0 321-60-8 1718-51-0	Nitrobenzene-d5 2-Fluorobiphenyl Terphenyl-d14	93% 92% 97%		24-125% 19-127% 10-119%	ó





MDL = Method Detection Limit



RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

By

XPL

n/a

Page 1 of 1

Client Sample ID: MW-16D Lab Sample ID: IC16038-6

Matrix: Method:

Project:

AQ - Ground Water SW846-8015C (DAI)

DF

1

BMSMC, Building 5 Area, PR

Analyzed

03/18/16

Date Sampled: 03/09/16 Date Received: 03/11/16

n/a

Q

Percent Solids:

n/a

Prep Date Prep Batch **Analytical Batch GGH5213**

Run #1 Run #2

Low Molecular Alcohol List

File ID

GH103826.D

CAS No.	Compound	Result	RL	MDL	Units
64-17-5	Ethanol	ND	100	55	ug/l
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l
67-56-1	Methanol	ND	200	71	ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limi	ts
111-27-3	Hexanol	98%		56-14	15%
111-27-3	Hexanol	94%		56-14	15%





MDL = Method Detection Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Report of Analysis

Page 1 of 2

Client Sample ID: TB030902 Lab Sample ID: JC16038-7

Matrix: AQ - Trip Blank Water

Method: SW846 8260C Project:

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16

Date Received: 03/11/16

Percent Solids: n/a

Q

Run #2

Purge Volume

Run #1 5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	ND	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/I
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	ND	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	ND	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	ND	1.0	0.27	ug/l
75-71 - 8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1, I-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	ND	1.0	0.27	ug/l
76-13-1	Freon 113	ND	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l

Méndez 10 = 188

ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 2

Client Sample ID: Lab Sample ID:

TB030902 JC16038-7

Matrix:

AQ - Trip Blank Water

Method: Project:

SW846 8260C

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a



VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	ND	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	ND	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	ND	1.0	0.38	ug/l	
95-47-6	o-Xylene	ND	1.0	0.17	ug/l	
1330-20-7	Xylene (total)	ND	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limi	ts	
1868-53-7	Dibromofluoromethane	105%		76-12	20%	
17060-07-0	1,2-Dichloroethane-D4	108%		73-12	22%	
2037-26-5	Toluene-D8	100%		84-11	19%	
460-00-4	4-Bromofluorobenzene	103%		78-11	17%	30
					98	30/



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Report of Analysis

Page 1 of 1

TB030902 Client Sample ID: Lab Sample ID: JC16038-7

Matrix: Method:

Project:

AQ - Trip Blank Water SW846-8015C (DAI)

BMSMC, Building 5 Area, PR

Date Sampled: 03/09/16 Date Received: 03/11/16

Percent Solids: n/a

Q

File ID DF Analyzed By Prep Date Prep Batch Run #1 GH103829.D 1 03/18/16 XPL n/a n/a Run #2	Analytical Batch GGH5213
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Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units
64-17-5	Ethanol	ND	100	55	ug/l
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l
67-63-0	Isopropyt Alcohol	ND	100	68	ug/l
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l
67-56-1	Methanol	ND	200	71	ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its
111-27-3	Hexanoi	99%		56-1	45%
111-27-3	Hexanol	91%		56-1	45%
					/





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Report of Analysis

By

NH

Prep Date

n/a

Page 1 of 2

Client Sample ID: S-30

JC16038-8 Lab Sample ID:

File ID

U204162.D

Matrix:

AQ - Ground Water SW846 8260C

Method: Project:

BMSMC, Building 5 Area, PR

Analyzed

03/15/16

DF

1

Date Sampled: 03/08/16 Date Received:

03/11/16

Percent Solids:

n/a

Q

Prep Batch **Analytical Batch**

VU9383

Run #1 Run #2

Purge Volume

Run #1 5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	ND	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	ND	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	ND	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	ND	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l \
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	ND	1.0	0.27	ug/l
76-13-1	Freon 113	ND	5.0	0.52	ug/l
591-78-6	2-Нехапопе	ND	5.0	1.7	ug/l



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 2

Client Sample ID: S-30

Lab Sample ID:JC16038-8Date Sampled:Matrix:AQ - Ground WaterDate Received:

Method: SW846 8260C

Project: BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16 Percent Solids: n/a



VOA TCL List

CAS No.	Compound	Result	RL.	MDL	Units	Q
98-82-8	Isopropylbenzene	ND	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	ND	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	ND	1.0	0.38	ug/l	
95-47-6	o-Xylene	ND	1.0	0.17	ug/l	
1330-20-7	Xylene (total)	ND	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limi	ts	
1868-53-7	Dibromofluoromethane	105%		76-13	20%	
17060-07-0	1,2-Dichloroethane-D4	107%		73-13	22%	
2037-26-5	Toluene-D8	99%		84-1	19%	
460-00-4	4-Bromofluorobenzene	104%		78-1	17%	09



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Page 1 of 3

Client Sample ID: Lab Sample ID:

S-30 JC16038-8

Matrix:

AQ - Ground Water SW846 8270D SW846 3510C Date Received:

Date Sampled:

Q

03/08/16 03/11/16

Method: Project:

BMSMC, Building 5 Area, PR

Percent Solids:

	====		
Prep Date	Prep Batch	Analytic	al Batch

File ID DF Analyzed By Run #1 P103411.D 1 03/17/16 SD 03/15/16 OP92078 EP4545 Run #2 P103375.D 40 03/16/16 LK 03/15/16 OP92078 EP4542

Initial Volume Final Volume

Run #1 990 ml Run #2 990 ml 1.0 ml 1.0 mi

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Units
95-57-8	2-Chlorophenol	ND	5.1	0.94	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.1	1.4	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.0	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.1	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	10	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.1	0.88	ug/l
95-48-7	2-Methylphenol	ND	2.0	0.83	ug/l
	3&4-Methylphenol	ND	2.0	0.68	ug/l
88-75-5	2-Nitrophenol	ND	5.1	1.4	ug/l
100-02-7	4-Nitrophenol	ND	10	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.1	1.5	ug/l
108-95-2	Phenol	ND	2.0	0.32	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.1	1.4	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.1	1.5	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.1	1.5	ug/l
83-32-9	Acenaphthene	ND	1.0	0.29	ug/l
208-96-8	Acenaphthylene	ND	1.0	0.24	ug/l
98-86-2	Acetophenone	ND	2.0	0.28	ug/l
120-12-7	Anthracene	ND	1.0	0.25	ug/l
1912-24-9	Atrazine	ND	2.0	0.42	ug/l
100-52-7	Benzaldehyde	0.90	5.1	0.34	ug/l
56-55-3	Benzo(a)anthracene	ND	1.0	0.32	ug/l
50-32-8	Benzo(a)pyrene	ND	1.0	0.34	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.0	0.32	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.0	0.41	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.0	0.37	ug/l /
101-55-3	4-Bromophenyl phenyl ether	ND	2.0	0.37	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.0	0.27	ug/l
92-52-4	1,1 -Biphenyl	ND	1.0	0.26	ug/l
91-58-7	2-Chloronaphthalene	ND	2.0	0.30	ug/l
106-47-8	4-Chloroaniline	ND	5.1	0.23	ug/l
86-74-8	Carbazole	ND	1.0	0.30	ug/l

tael Infant Méndez 16 # 1886

ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 3

Client Sample ID: S-30

Lab Sample ID: JC16038-8

Matrix: AQ - Ground Water Method: SW846 8270D SW8

Method: SW846 8270D SW846 3510C Project: BMSMC, Building 5 Area, PR Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids: n/a

Q



ABN TCL List (SOM0 1.1)

	•				
CAS No.	Compound	Result	RL	MDL	Units
105-60-2	Caprolactam	ND	2.0	0.43	ug/l
218-01-9	Chrysene	ND	1.0	0.35	ug/l
111-91-1	bis (2-Chloroethoxy) methane	ND	2.0	0.26	ug/l
111-44-4	bis(2-Chloroethyl)ether	ND	2.0	0.35	ug/l
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.0	0.29	ug/l
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.0	0.27	ug/l
121-14-2	2,4-Dinitrotoluene	ND	1.0	0.27	ug/l
606-20-2	2,6-Dinitrotoluene	ND	1.0	0.33	ug/l
91-94-1	3,3'-Dichlorobenzidine	ND	2.0	0.54	ug/l
123-91-1	1,4-Dioxane	2220 ª	40	29	ug/l
53-70-3	Dibenzo(a,h)anthracene	ND	1.0	0.37	ug/l
132-64-9	Dibenzofuran	ND	5.1	0.27	ug/l
84-74-2	Di-n-butyl phthalate	ND	2.0	0.79	ug/l
117-84-0	Di-n-octyl phthalate	ND	2.0	0.29	ug/l
84-66-2	Diethyl phthalate	ND	2.0	0.25	ug/l
131-11-3	Dimethyl phthalate	ND	2.0	0.32	ug/l
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.0	0.78	ug/l
206-44-0	Fluoranthene	ND	1.0	0.23	ug/l
86-73-7	Fluorene	ND	1.0	0.30	ug/l
118-74-1	Hexachlorobenzene	ND	1.0	0.43	ug/l
87-68-3	Hexachlorobutadiene	ND	1.0	0.37	ug/l
77-47-4	Hexachlorocyclopentadiene	ND	10	0.30	ug/l
67-72-1	Hexachloroethane	ND	2.0	0.22	ug/l
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.0	0.39	ug/l
78-59-1	Isophorone	ND	2.0	0.29	ug/l
91-57-6	2-Methylnaphthalene	ND	1.0	0.29	ug/l
88-74-4	2-Nitroaniline	ND	5.1	0.21	ug/l
99-09-2	3-Nitroaniline	ND	5.1	0.24	ug/l
100-01-6	4-Nitroaniline	ND	5.1	0.35	ug/l
91-20-3	Naphthalene	ND	1.0	0.29	ug/l
98-95-3	Nitrobenzene	ND	2.0	0.47	ug/l
621-64-7	N-Nitroso-di-n-propylamine	ND	2.0	0.32	ug/l
86-30-6	N-Nitrosodiphenylamine	ND	5.1	0.30	ug/l
85-01-8	Phenanthrene	ND	1.0	0.23	ug/l
129-00-0	Pvrene	ND	1.0	0.34	ug/l
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.0	0.37	ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its

67%

61%



RL = Reporting Limit

367-12-4

2-Fluorophenol

14-88%

fuel Infante Mendez

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Method:

Project:

Report of Analysis

Page 3 of 3

Client Sample ID: S-30 Lab Sample ID:

JC16038-8 Matrix: AQ - Ground Water

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16 Percent Solids: n/a

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limits
4165-62-2	Phenol-d5	45%	34%	10-110%
118-79-6	2,4,6-Tribromophenol	105%	55%	39-149%
4165-60-0	Nitrobenzene-d5	101%	93%	32-128%
321-60-8	2-Fluorobiphenyl	95%	120% և	35-119%
1718-51-0	Terphenyl-d14	96%	84%	10-126%

(a) Result is from Run# 2

(b) Outside control limits due to dilution.





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 1 of 1

Client Sample ID: S-30

Lab Sample ID: JC16038-8

Matrix:

AQ - Ground Water

Method:

SW846 8270D BY SIM SW846 3510C

Project:

BMSMC, Building 5 Area, PR

Date Sampled:

03/08/16 Date Received: 03/11/16

Percent Solids:

File ID DF Analyzed Prep Date Prep Batch **Analytical Batch** By Run #1 4P15642.D 1 03/17/16 IJ 03/15/16 OP92078A E4P804

Run #2

Initial Volume Final Volume Run #1 990 ml 1.0 ml

Run #2

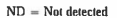
CAS No. Compound Result RL MDL Units Q

91-20-3 Naphthalene ND 0.10 0.013 ug/l

CAS No. Surrogate Recoveries Run#1 Run# 2 Limits

4165-60-0 Nitrobenzene-d5 94% 24-125% 321-60-8 2-Fluorobiphenyl 96% 19-127% 1718-51-0 Terphenyl-d14 93% 10-119%





MDL = Method Detection Limit



E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Report of Analysis

Page 1 of 1

S-30 Client Sample ID:

Lab Sample ID: JC16038-8

Matrix:

AQ - Ground Water

Method: Project:

SW846-8015C (DAI)

BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids:

T3 21 4	File ID	DF	Analyzed	Ву	Prep Date	Prep Batch	Analytical Batch
Run #1 Run #2	GH103789.D	1	03/17/16	XPL	n/a	n/a	GGH5211

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units	Q
64-17-5	Ethanol	ND	100	55	ug/l	
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l	
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l	
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l	
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l	
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l	
67-56-1	Methanol	ND	200	71	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
111-27-3	Hexanol	98%		56-1	45%	, el
111-27-3	Hexanol	90%			45%	ak la





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Report of Analysis

By

BP

0.010

Prep Date

03/15/16

MDL

0.0042

0.0049

Units

ug/l

ug/l

Page 1 of 1

Client Sample ID: S-30

JC16038-8 Lab Sample ID:

File ID

4G66288.D

4,4'-DDD

Matrix:

AQ - Ground Water

DF

1

Method: Project:

SW846 8081B SW846 3510C

BMSMC, Building 5 Area, PR

Analyzed

03/20/16

ND

Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids:

Prep Batch

OP92108

Q

Analytical Batch G4G1744

Run #1 Run #2

Initial Volume Final Volume Run #1 1000 ml $10.0 \, ml$

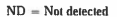
Run #2

72-54-8

CAS No. Compound Result RL 319-85-7 beta-BHC ND 0.010

50-29-3	4,4'-DDT	ND	0.010	0.0047 ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limits
877-D9-8	Tetrachloro-m-xylene	97%		26-132%
877-09-8	Tetrachloro-m-xylene	99%		26-132%
2051-24-3	Decachlorobiphenyl	101%		10-118%
2051-24-3	Decachlorobiphenyl	103%		10-118%





MDL = Method Detection Limit



RL = Reporting Limit

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B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Report of Analysis

Page 1 of 2

Client Sample ID:

FB030816

Lab Sample ID: JC16038-9

Matrix:

AQ - Field Blank Water

SW846 8260C

Method: Project:

BMSMC, Building 5 Area, PR

Date Sampled:

Q

J

03/08/16

Date Received:

03/11/16

Percent Solids:

File ID DF Analyzed By Prep Date Prep Batch **Analytical Batch** Run #1 U204160.D 03/15/16 NH n/a n/a VU9383 Run #2

Purge Volume

Run #1

5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Unit
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	0.30	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	ND	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	ND	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	ND	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1,1-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	ND	1.0	0.27	ug/l
76-13-1	Freon 113	ND	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l
					-



ND = Not detected

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Page 2 of 2

Client Sample ID:

FB030816

Lab Sample ID: Matrix:

Method:

Project:

JC16038-9

AQ - Field Blank Water

SW846 8260C

BMSMC, Building 5 Area, PR

Date Sampled: 03
Date Received: 03

03/08/16 03/11/16

Percent Solids: n/a



VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	0.91	1.0	0.23	ug/l	J
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	-
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	4.5	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachioroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	ND	1.0	0.38	ug/l	
95-47-6	o-Xylene	ND	1.0	0.17	ug/l	
1330-20-7	Xylene (total)	ND	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its	
1868-53-7	Dibromofluoromethane	105%		76-1	20%	
17060-07-0	1,2-Dichloroethane-D4	107%		73-1	22%	
2037-26-5	Toluene-D8	99%		84-1	19%	
460-00-4	4-Bromofluorobenzene	103%		78-1	17%	



ND = Not detected

MDL = Method Detection Limit

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B = Indicates analyte found in associated method blank

Ву

SD

Prep Date

03/15/16

Page 1 of 3

Client Sample ID: FB030816 Lab Sample ID: JC16038-9

File ID

F155576.D

AQ - Field Blank Water

Date Sampled: 03/08/16 Date Received: 03/11/16

Matrix: Method:

SW846 8270D SW846 3510C

Percent Solids: n/a

Q

Project:

BMSMC, Building 5 Area, PR

Analyzed

03/15/16

Prep Batch OP92078

Analytical Batch EF6543

Run #1 Run #2

Initial Volume Final Volume 1000 ml

Run #1

1.0 ml

DF

1

Run #2

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Units
95-57-8	2-Chlorophenol	ND	5.0	0.93	ug/l
59-50-7	4-Chloro-3-methyl phenol	ND	5.0	1.4	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.0	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.0	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	10	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.0	0.87	ug/l
95-48-7	2-Methylphenol	ND	2.0	0.82	ug/l
	3&4-Methylphenol	ND	2.0	0.67	ug/l
88-75-5	2-Nitrophenol	ND	5.0	1.4	ug/l
100-02-7	4-Nitrophenol	ND	10	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.0	1.4	ug/l
108-95-2	Phenol	ND	2.0	0.31	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.0	1.4	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.0	1.5	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.0	1.4	ug/l
83-32-9	Acenaphthene	ND	1.0	0.29	ug/l
208-96-8	Acenaphthylene	ND	1.0	0.24	ug/l
98-86-2	Acetophenone	ND	2.0	0.28	ug/l
120-12-7	Anthracene	ND	1.0	0.25	ug/l
1912-24-9	Atrazine	ND	2.0	0.42	ug/l
100-52-7	Benzaldehyde	ND	5.0	0.34	ug/l
56-55-3	Benzo(a)anthracene	ND	1.0	0.32	ug/l
50-32-8	Benzo(a)pyrene	ND	1.0	0.33	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.0	0.32	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.0	0.41	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.0	0.37	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.0	0.37	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.0	0.27	ug/l
92-52-4	1,1 -Biphenyl	ND	1.0	0.26	ug/l
91-58-7	2-Chloronaphthalene	ND	2.0	0.30	ug/l
106-47-8	4-Chloroaniline	ND	5.0	0.23	ug/l
86-74-8	Carbazole	ND	1.0	0.29	ug/l



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Project:

Report of Analysis

Page 2 of 3

FB030816 Client Sample ID: Lab Sample ID:

JC16038-9 Matrix: AQ - Field Blank Water Method:

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16 n/a

Percent Solids:



ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Units	Q	
105-60-2	Caprolactam	ND	2.0	0.43	ug/l		
218-01-9	Chrysene	ND	1.0	0.35	ug/l		
111-91-1	bis(2-Chloroethoxy)methane	ND	2.0	0.26	ug/l		
111-44-4	bis(2-Chloroethyl)ether	ND	2.0	0.34	ug/l		
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.0	0.28	ug/l		
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.0	0.27	ug/l		
121-14-2	2,4-Dinitrotoluene	ND	1.0	0.26	ug/l		
606-20-2	2,6-Dinitrotoluene	ND	1.0	0.32	ug/l		
91-94-1	3,3'-Dichlorobenzidine	ND	2.0	0.53	ug/l		
53-70-3	Dibenzo(a,h)anthracene	ND	1.0	0.37	ug/l		
132-64-9	Dibenzofuran	ND	5.0	0.27	ug/l		
84-74-2	Di-n-butyl phthalate	ND	2.0	0.79	ug/l		
117-84-0	Di-n-octyl phthalate	ND	2.0	0.29	ug/l		
84-66-2	Diethyl phthalate	ND	2.0	0.24	ug/l		
131-11-3	Dimethyl phthalate	ND	2.0	0.31	ug/l		
117-81-7	bis(2-Ethylhexyl)phthalate	1.2	2.0	0.77	ug/l	J	
206-44-0	Fluoranthene	ND	1.0	0.23	ug/l		
86-73-7	Fluorene	ND	1.0	0.29	ug/l		
118-74-1	Hexachlorobenzene	ND	1.0	0.42	ug/l		
87-68-3	Hexachlorobutadiene	ND	1.0	0.36	ug/l		
77-47-4	Hexachlorocyclopentadiene	ND	10	0.29	ug/l		
67-72-1	Hexachloroethane	ND	2.0	0.22	ug/l		
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.0	0.38	ug/l		
78-59-1	Isophorone	ND	2.0	0.29	ug/l		
91-57-6	2-Methylnaphthalene	ND	1.0	0.29	ug/l		
88-74-4	2-Nitroaniline	ND	5.0	0.21	ug/l		
99-09-2	3-Nitroaniline	ND	5.0	0.24	ug/l		
100-01-6	4-Nitroaniline	ND	5.0	0.34	ug/l		
91-20-3	Naphthalene	ND	1.0	0.28	ug/l		
98-95-3	Nitrobenzene	ND	2.0	0.46	ug/l		
621-64-7	N-Nitroso-di-n-propylamine	ND	2.0	0.31	ug/l		
86-30-6	N-Nitrosodiphenylamine	ND	5.0	0.29	ug/l		
85-01-8	Phenanthrene	ND	1.0	0.23	ug/l	1	
129-00-0	Pyrene	ND	1.0	0.34	ug/l	- (
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.0	0.36	ug/l	1	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its	\	
367-12-4	2-Fluorophenol	51%		14-8	8%		
4165-62-2	Phenol-d5	36%		10-1	10%		

ND = Not detected MDL = Method Detection Limit

J = Indicates an estimated value

RL = Reporting Limit E = Indicates value exceeds calibration range B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

fael Infante Méndez

Report of Analysis

Page 3 of 3

Client Sample ID: FB030816 Lab Sample ID: JC16038-9

AQ - Field Blank Water

Date Sampled: Date Received:

03/08/16 03/11/16 Percent Solids:

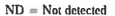
Matrix: Method: Project:

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limits
118-79-6	2,4,6-Tribromophenol	87%		39-149%
4165-60-0	Nitrobenzene-d5	90%		32-128%
321-60-8	2-Fluorobiphenyl	83%		35-119%
1718-51-0	Terphenyl-d14	83%		10-126%





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Client Sample ID: FB030816 Lab Sample ID: JC16038-9

Matrix: AQ - Field Blank Water

Method: SW846 8270D BY SIM SW846 3510C

Project: BMSMC, Building 5 Area, PR Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids:

File ID DF Analyzed Ву Prep Date Prep Batch **Analytical Batch** Run #1 4P15643.D 1 03/17/16 11 03/15/16 OP92078A E4P804

Run #2

Initial Volume Final Volume Run #1 1000 ml 1.0 ml

Run #2

CAS No.	Compound	Result	RL	MDL	Units	Q
91-20-3 123-91-1	Naphthalene 1,4-Dioxane	ND ND	0.10 0.10	0.013 0.053	ug/l ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its	
4165-60-0 321-60-8 1718-51-0	Nitrobenzene-d5 2-Fluorobiphenyl Terphenyl-d14	90% 96% 92%		19-1	25% 27% 19%	





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

Report of Analysis

Page 1 of 1

Client Sample ID: FB030816 Lab Sample ID:

JC16038-9

AQ - Field Blank Water SW846-8015C (DAI)

Date Sampled: 03/08/16 Date Received: 03/11/16

Q

Percent Solids: n/a

Method: Project:

Matrix:

BMSMC, Building 5 Area, PR

File ID DF Analyzed Ву Prep Date Prep Batch **Analytical Batch** Run #1 GH103799.D 03/17/16 XPL n/a n/a **GGH5211**

Run #2

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units
64-17-5	Ethanol	ND	100	55	ug/l
78-83-1	Isobutyl Alcohol	ND	100	36	ug/l
67-63-0	Isopropyl Alcohol	ND	100	68	ug/l
71-23-8	n-Propyl Alcohol	ND	100	43	ug/l
71-36-3	n-Butyl Alcohol	ND	100	87	ug/l
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l
67-56-1	Methanol	ND	200	71	ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Lim	its
111-27-3	Hexanol	102%		56-1	45%
111-27-3	Hexanol	90%		56-1	45%





MDL = Method Detection Limit

RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



E = Indicates value exceeds calibration range

Matrix:

Method:

Project:

Report of Analysis

Page 1 of 1

Client Sample ID: FB030816 Lab Sample ID: JC16038-9

AQ - Field Blank Water SW846 8081B SW846 3510C BMSMC, Building 5 Area, PR

03/08/16 Date Sampled: Date Received: 03/11/16

Percent Solids:

Q

File ID **Analytical Batch** DF Analyzed By Prep Date Prep Batch Run #1 4G66287.D 1 03/20/16 BP 03/15/16 OP92108 G4G1744 Run #2

Initial Volume Final Volume Run #1 1000 ml 10.0 ml

Run #2

CAS No.	Compound	Result	RL	MDL	Units
319-85-7 72-54-8 50-29-3	heta-BHC 4,4'-DDD 4,4'-DDT	ND ND ND	0.010 0.010 0.010	0.0042 0.0049 0.0047	ug/l ug/l ug/l
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limi	ts
877-09-8 877-09-8 2051-24-3 2051-24-3	Tetrachloro-m-xylene Tetrachloro-m-xylene Decachlorobiphenyl Decachlorobiphenyl	78% 86% 67% 73%		26-13 26-13 10-13	32% 18%



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Ву

NH

Prep Date

n/a

Page 1 of 2

Client Sample ID: MW-11

Lab Sample ID: JC16038-10

File ID

U204163.D

Matrix: Method: AQ - Ground Water

DF

1

SW846 8260C

Date Sampled: Date Received:

03/08/16 03/11/16

Percent Solids: n/a

Project:

BMSMC, Building 5 Area, PR

Analyzed

03/16/16

Prep Batch

Q

Analytical Batch

VU9383 n/a

Run #1 Run #2

Purge Volume

Run #1 5.0 ml

Run #2

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units
67-64-1	Acetone	ND	10	3.3	ug/l
71-43-2	Benzene	ND	0.50	0.24	ug/l
74-97-5	Bromochloromethane	ND	1.0	0.37	ug/l
75-27-4	Bromodichloromethane	ND	1.0	0.23	ug/l
75-25-2	Bromoform	ND	1.0	0.23	ug/l
74-83-9	Bromomethane	ND	2.0	0.42	ug/l
78-93-3	2-Butanone (MEK)	ND	10	5.6	ug/l
75-15-0	Carbon disulfide	ND	2.0	0.25	ug/l
56-23-5	Carbon tetrachloride	ND	1.0	0.22	ug/l
108-90-7	Chlorobenzene	ND	1.0	0.19	ug/l
75-00-3	Chloroethane	ND	1.0	0.34	ug/l
67-66-3	Chloroform	ND	1.0	0.19	ug/l
74-87-3	Chloromethane	ND	1.0	0.41	ug/l
110-82-7	Cyclohexane	ND	5.0	0.28	ug/l
96-12-8	1,2-Dibromo-3-chloropropane	ND	2.0	0.99	ug/l
124-48-1	Dibromochloromethane	ND	1.0	0.15	ug/l
106-93-4	1,2-Dibromoethane	ND	1.0	0.23	ug/l
95-50-1	1,2-Dichlorobenzene	ND	1.0	0.19	ug/l
541-73-1	1,3-Dichlorobenzene	ND	1.0	0.23	ug/l
106-46-7	1,4-Dichlorobenzene	ND	1.0	0.27	ug/l
75-71-8	Dichlorodifluoromethane	ND	2.0	0.90	ug/l
75-34-3	1,1-Dichloroethane	ND	1.0	0.17	ug/l
107-06-2	1,2-Dichloroethane	ND	1.0	0.18	ug/l
75-35-4	1, I-Dichloroethene	ND	1.0	0.51	ug/l
156-59-2	cis-1,2-Dichloroethene	ND	1.0	0.27	ug/l
156-60-5	trans-1,2-Dichloroethene	ND	1.0	0.65	ug/l
78-87-5	1,2-Dichloropropane	ND	1.0	0.39	ug/l
10061-01-5	cis-1,3-Dichloropropene	ND	1.0	0.21	ug/l
10061-02-6	trans-1,3-Dichloropropene	ND	1.0	0.19	ug/l
100-41-4	Ethylbenzene	ND	1.0	0.27	ug/l
76-13-1	Freon 113	ND	5.0	0.52	ug/l
591-78-6	2-Hexanone	ND	5.0	1.7	ug/l



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 2 of 2

Client Sample ID: MW-11 Lab Sample ID:

JC16038-10

Date Sampled:

03/08/16

Matrix:

AQ - Ground Water SW846 8260C

Date Received: Percent Solids: n/a

03/11/16

Method: Project:

BMSMC, Building 5 Area, PR

VOA TCL List

CAS No.	Compound	Result	RL	MDL	Units	Q
98-82-8	Isopropylbenzene	ND	1.0	0.23	ug/l	
79-20-9	Methyl Acetate	ND	5.0	1.9	ug/l	
108-87-2	Methylcyclohexane	ND	5.0	0.22	ug/l	
1634-04-4	Methyl Tert Butyl Ether	1.9	1.0	0.24	ug/l	
108-10-1	4-Methyl-2-pentanone(MIBK)	ND	5.0	1.0	ug/l	
75-09-2	Methylene chloride	ND	2.0	0.73	ug/l	
100-42-5	Styrene	ND	1.0	0.27	ug/l	
79-34-5	1,1,2,2-Tetrachloroethane	ND	1.0	0.21	ug/l	
127-18-4	Tetrachloroethene	ND	1.0	0.40	ug/l	
108-88-3	Toluene	ND	1.0	0.16	ug/l	
87-61-6	1,2,3-Trichlorobenzene	ND	1.0	0.23	ug/l	
120-82-1	1,2,4-Trichlorobenzene	ND	1.0	0.21	ug/l	
71-55-6	1,1,1-Trichloroethane	ND	1.0	0.25	ug/l	
79-00-5	1,1,2-Trichloroethane	ND	1.0	0.21	ug/l	
79-01-6	Trichloroethene	ND	1.0	0.22	ug/l	
75-69-4	Trichlorofluoromethane	ND	2.0	0.43	ug/l	
75-01-4	Vinyl chloride	ND	1.0	0.15	ug/l	
	m,p-Xylene	ND	1.0	0.38	ug/l	
95-47-6	o-Xylene	ND	1.0	0.17	ug/l	
1330-20-7	Xylene (total)	ND	1.0	0.17	ug/l	
CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limi	ts	
1868-53-7	Dibromofluoromethane	104%		76-17	20%	
17060-07-0	1,2-Dichloroethane-D4	107%		73-13	22%	
2037-26-5	Toluene-D8	99%		84-11	19%	
460-00-4	4-Bromofluorobenzene	103%		78-1	17%	





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

By

LK

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Page 1 of 3

Client Sample ID: MW-11

Lab Sample ID: JC16038-10

Matrix:

AQ - Ground Water

Date Sampled: 03 Date Received: 03

03/08/16 03/11/16

Method:

SW846 8270D SW846 3510C

Percent Solids: n/

n/a

Project:

BMSMC, Building 5 Area, PR

Q

Prep Batch Analytical Batch

File ID
Run #1 P103372.D
Run #2 P103374.D

DF Analyzed
1 03/16/16
5 03/16/16

Prep Date 03/15/16 03/15/16

OP92078 OP92078

EP4542 EP4542

Initial Volume Final Volume

Run #1 970 ml Run #2 970 ml 1.0 ml 1.0 ml

ABN TCL List (SOM0 1.1)

CAS No.	Compound	Result	RL	MDL	Units
95-57-8	2-Chlorophenol	ND	5.2	0.96	ug/l
59-50-7	4-Chloro-3-methyl phenoi	ND	5.2	1.4	ug/l
120-83-2	2,4-Dichlorophenol	ND	2.1	1.3	ug/l
105-67-9	2,4-Dimethylphenol	ND	5.2	1.3	ug/l
51-28-5	2,4-Dinitrophenol	ND	10	1.1	ug/l
534-52-1	4,6-Dinitro-o-cresol	ND	5.2	0.90	ug/l
95-48-7	2-Methylphenol	ND	2.1	0.84	ug/l
	3&4-Methylphenol	ND	2.1	0.69	ug/l
88-75-5	2-Nitrophenol	ND	5.2	1.5	ug/l
100-02-7	4-Nitrophenol	ND	10	1.1	ug/l
87-86-5	Pentachlorophenol	ND	5.2	1.5	ug/l
108-95-2	Phenol	ND	2.1	0.32	ug/l
58-90-2	2,3,4,6-Tetrachlorophenol	ND	5.2	1.5	ug/l
95-95-4	2,4,5-Trichlorophenol	ND	5.2	1.5	ug/l
88-06-2	2,4,6-Trichlorophenol	ND	5.2	1.5	ug/l
83-32-9	Acenaphthene	ND	1.0	0.29	ug/l
208-96-8	Acenaphthylene	ND	1.0	0.25	ug/l
98-86-2	Acetophenone	ND	2.1	0.28	ug/l
120-12-7	Anthracene	ND	1.0	0.25	ug/l
1912-24-9	Atrazine	ND	2.1	0.43	ug/l
100-52-7	Benzaldehyde	ND	5.2	0.35	ug/l
56-55-3	Benzo(a)anthracene	ND	1.0	0.32	ug/l
50-32-8	Вепго(а)ругене	ND	1.0	0.34	ug/l
205-99-2	Benzo(b)fluoranthene	ND	1.0	0.33	ug/l
191-24-2	Benzo(g,h,i)perylene	ND	1.0	0.42	ug/l
207-08-9	Benzo(k)fluoranthene	ND	1.0	0.38	ug/l
101-55-3	4-Bromophenyl phenyl ether	ND	2.1	0.38	ug/l
85-68-7	Butyl benzyl phthalate	ND	2.1	0.28	ug/l
92-52-4	1,1'-Biphenyl	ND	1.0	0.26	ug/l
91-58-7	2-Chloronaphthalene	ND	2.1	0.31	ug/l
106-47-8	4-Chloroaniline	ND	5.2	0.24	ug/l
86-74-8	Carbazole	ND	1.0	0.30	ug/l



ND = Not detected

MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

Page 2 of 3

Client Sample ID: MW-11

Lab Sample ID: JC16038-10

Matrix: Method:

Project:

AQ - Ground Water

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR Date Sampled: 0
Date Received: 0

03/08/16 03/11/16

Percent Solids: n/a

ABN TCL List (SOM0 1.1)

	Bas (BO145 1:1)					
CAS No.	Compound	Result	RL	MDL	Units	Q
105-60-2	Caprolactam	ND	2.1	0.44	ug/l	
218-01-9	Chrysene	ND	1.0	0.36	ug/l	
111-91-1	bis(2-Chloroethoxy)methane	ND	2.1	0.27	ug/l	
111-44-4	bis(2-Chloroethyl)ether	ND	2.1	0.35	ug/l	
108-60-1	bis(2-Chloroisopropyl)ether	ND	2.1	0.29	ug/l	
7005-72-3	4-Chlorophenyl phenyl ether	ND	2.1	0.28	ug/l	
121-14-2	2,4-Dinitrotoluene	ND	1.0	0.27	ug/l	
606-20-2	2,6-Dinitrotoluene	ND	1.0	0.33	ug/l	
91-94-1	3,3'-Dichlorobenzidine	ND	2.1	0.55	ug/l	
123-91-1	1,4-Dioxane	385 a	5.2	3.7	ug/l	
53-70-3	Dibenzo(a,h)anthracene	ND	1.0	0.38	ug/l	
132-64-9	Dibenzofuran	ND	5.2	0.28	ug/l	
84-74-2	Di-n-butyl phthalate	ND	2.1	0.81	ug/l	
117-84-0	Di-n-octyl phthalate	ND	2.1	0.30	ug/l	
84-66-2	Diethyl phthalate	ND	2.1	0.25	ug/l	
131-11-3	Dimethyl phthalate	ND	2.1	0.32	ug/l	
117-81-7	bis(2-Ethylhexyl)phthalate	ND	2.1	0.79	ug/l	
206-44-0	Fluoranthene	ND	1.0	0.24	ug/l	
86-73-7	Fluorene	ND	1.0	0.30	ug/l	
118-74-1	Hexachlorobenzene	ND	1.0	0.44	ug/l	
87-68-3	Hexachlorobutadiene	ND	1.0	0.38	ug/l	
77-47-4	Hexachlorocyclopentadiene	ND	10	0.30	ug/l	
67-72-1	Hexachloroethane	ND	2.1	0.23	ug/l	
193-39-5	Indeno(1,2,3-cd)pyrene	ND	1.0	0.40	ug/l	
78-59-1	Isophorone	ND	2.1	0.29	ug/l	
91-57-6	2-Methylnaphthalene	ND	1.0	0.30	ug/l	
88-74-4	2-Nitroaniline	ND	5.2	0.22	ug/l	
99-09-2	3-Nitroaniline	ND	5.2	0.25	ug/l	
100-01-6	4-Nitroaniline	ND	5.2	0.35	ug/l	
91-20-3	Naphthalene	ND	1.0	0.29	ug/l	
98-95-3	Nitrobenzene	ND	2.1	0.48	ug/l	
621-64-7	N-Nitroso-di-n-propylamine	ND	2.1	0.32	ug/l	CONTINUE
86-30-6	N-Nitrosodiphenylamine	ND	5.2	0.30	ug/l	332
85-01-8	Phenanthrene	ND	1.0	0.24	ug/l	fael Infa
129-00-0	Pyrene	ND	1.0	0.35	ug/l	HIGH HIME
95-94-3	1,2,4,5-Tetrachlorobenzene	ND	2.1	0.37	ug/l	IC 18
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	Upper
367-12-4	2-Fluorophenol	65%	63%	14-8	8%	SEO FICE



RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank

E = Indicates value exceeds calibration range

Page 3 of 3

MW-11 Client Sample ID:

Lab Sample ID: JC16038-10

Matrix: Method: Project:

AQ - Ground Water

SW846 8270D SW846 3510C BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids:

ABN TCL List (SOM0 1.1)

CAS No.	Surrogate Recoveries	Run#1	Run# 2	Limits
4165-62-2	Phenol-d5	46%	46%	10-110%
118-79-6	2,4,6-Tribromophenol	99%	90%	39-149%
4165-60-0	Nitrobenzene-d5	97%	104%	32-128%
321-60-8	2-Fluorobiphenyl	95%	113%	35-119%
1718-51-0	Terphenyl-d14	93%	99%	10-126%

(a) Result is from Run# 2





MDL = Method Detection Limit

RL = Reporting Limit

E = Indicates value exceeds calibration range

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



Page 1 of 1

MW-11 Client Sample ID:

Lab Sample ID: JC16038-10 Matrix:

Method: Project:

AQ - Ground Water SW846 8270D BY SIM SW846 3510C

BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids: n/a

Prep Date File ID DF Analyzed By Prep Batch **Analytical Batch** 4P15644.D 03/15/16 OP92078A E4P804 Run #1 1 03/17/16 IJ

Run #2

Initial Volume Final Volume Run #1 970 ml

Run #2

4165-60-0

321-60-8

1718-51-0

1.0 ml

CAS No. Compound Result

RL

Run#2

Units

Q

91-20-3 Naphthalene ND

Run#1

0.10

0.014 ug/l

MDL

CAS No. Surrogate Recoveries

Nitrobenzene-d5

2-Fluorobiphenyl

Terphenyl-d14

110% 115% 24-125% 19-127%

10-119%

Limits

111%

MDL = Method Detection Limit





RL = Reporting Limit E = Indicates value exceeds calibration range



J = Indicates an estimated value

B = Indicates analyte found in associated method blank

N = Indicates presumptive evidence of a compound

Method:

Project:

Report of Analysis

Page 1 of 1

Client Sample ID: MW-11

Lab Sample ID: JC16038-10 Matrix:

AQ - Ground Water SW846-8015C (DAI)

BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids: n/a

	File ID	DF	Analyzed	By	Prep Date	Prep Batch	Analytical Batch
Run #1	GH103788.D	1	03/17/16	XPL	n/a	n/a	GGH5211
Run #2							!

Low Molecular Alcohol List

CAS No.	Compound	Result	RL	MDL	Units	Q
64-17-5	Ethanol	ND	100	55	ug/l	
78-83-1 67-63-0	Isobutyl Alcohol Isopropyl Alcohol	ND ND	100 100	36 68	ug/l ug/l	
71-23-8 71-36-3	n-Propyl Alcohol n-Butyl Alcohol	ND ND	100 100	43 87	ug/l	
78-92-2	sec-Butyl Alcohol	ND	100	66	ug/l ug/l	
67-56-1	Methanol	ND	200	71	ug/l	
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Lim	its	
111-27-3	Hexanol	95%		56-1	45%	
111-27-3	Hexanol	85%		56-1	45%	





MDL = Method Detection Limit

RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



E = Indicates value exceeds calibration range

Report of Analysis

Page 1 of 1

Client Sample ID: MW-11 Lab Sample ID: JC16038-10

Matrix:

AQ - Ground Water SW846 8081B SW846 3510C

BMSMC, Building 5 Area, PR

Date Sampled: 03/08/16 Date Received: 03/11/16

Percent Solids: n/a

Q

File ID DF Analyzed By Prep Date Prep Batch **Analytical Batch** Run #1 4G66289.D 03/20/16 BP G4G1744 1 03/15/16 OP92108

Run #2

Method:

Project:

Initial Volume Final Volume Run #1 940 ml 10.0 ml

Run #2

CAS No.	Compound	Result	RL	MDL	Units
319-85-7 72-54-8 50-29-3	beta-BHC 4,4'-DDD 4,4'-DDT	ND ND ND	0.011 0.011 0.011	0.0045 0.0052 0.0050	ug/l ug/l ug/l
CAS No.	Surrogate Recoveries	Run# 1	Run# 2	Limi	ts
877-09-8 877-09-8 2051-24-3 2051-24-3	Tetrachloro-m-xylene Tetrachloro-m-xylene Decachlorobiphenyl Decachlorobiphenyl	69% 76% 72% 81%		26-13 26-13 10-11 10-11	2% 8%





MDL = Method Detection Limit

RL = Reporting Limit

J = Indicates an estimated value

B = Indicates analyte found in associated method blank



E = Indicates value exceeds calibration range

															Accustment Job #: JC/16038 Accustment Quadra II:							
Client Information Facility Information												Anal	rtical Infe	rmation								
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JC16038: Chain of Custody Page 1 of 4

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JC16038: Chain of Custody Page 2 of 4

EXECUTIVE NARRATIVE

SDG No:

JC16038

Laboratory:

Accutest, New Jersey

Analysis:

SW846-8260C

Number of Samples:

10

Location:

BMSMC, Building 5 Area

Humacao, PR

SUMMARY:

Eight (8) groundwater samples, one field blank, and one trip blank were analyzed for the VOA TCL list following method SW846-8260C. The sample results were assessed according to USEPA data validation guidance documents in the following order of precedence Hazardous Waste Support Section SOP No. HW-33A, Revision 0, June, 2015. SOM02.2. Low/Medium Volatile Data Validation. The QC criteria and data validation actions listed on the data review worksheets are from the primary guidance document, unless otherwise noted.

Results are valid and can be used for decision making purposes.

Critical issues:

None

Major:

None

Minor:

1. Initial calibration and initial calibration verification within the required criteria. Continuing calibration for Freon 113 (- 28% D) and Acetone (-33.3 % D) outside method performance criteria but within guidance document criteria. Closing calibration check verification not included in data package. No action

taken, professional judgment.

2. Target analytes (chlorobenzene, isopropyl benzene, and MTBE) fould in field

blank. No action taken.

3. MS/MSD % recoveries outside laboratory control limits for sample JC16250-14MS/-14MSD. No action taken, sample concentration high compared to

amount spiked.

Critical findings:

None

Major findings:

None

Minor findings:

None

COMMENTS:

Results are valid and can be used for decision making purposes.

Reviewers Name:

Rafael Infante

Chemist License 1889

Signature:

April 16, 2016

Date:

SAMPLE ORGANIC DATA SAMPLE SUMMARY

Sample ID: JC16038-1

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units E	ilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	0.5	ug/L	1.0	-	U	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	0.28	ug/L	1.0	J	UJ	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	-	U	Yes
Cyclohexane	2.0	ug/L	1.0	-	U	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	0.90	ug/L	1.0	J	UJ	Yes
1,3-Dichlorobenzene	1.0	ug/L	1.0	•	U	Yes
1,4-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
trans-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloropropane	1.0	ug/L	1.0	-	U	Yes
cis-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes
trans-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes
Ethylbenzene	1.0	ug/L	1.0	-	U	Yes
Freon 113	1.0	ug/L	1.0	-	U	Yes
2-Hexanone	5.0	ug/L	1.0	-	U	Yes
Isopropylbenzene	1.0	ug/L	1.0	-	U	Yes
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes
Methylcyclohexane	5.0	ug/L	1.0	-	U	Yes
Methyl Tert Butyl Ether	1.0	ug/L	1.0	-	U	Yes
4-Methyl-2-pentanone(MIBK)	2.0	ug/L	1.0	-	U	Yes
Methylene chloride	2.0	ug/L	1.0	-	U	Yes
Styrene	1.0	ug/L	1.0	-	U	Yes
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0	-	U	Yes
Tetrachloroethene	1.0	ug/L	1.0	-	U	Yes
Tetrahydrofuran	10	ug/L	1.0	-	U	Yes
Toluene	1.0	ug/L	1.0	-	U	Yes
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,1,1-Trichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1,2-Trichloroethane	1.0	ug/L	1.0	-	U	Yes
Trichloroethene	1.0	ug/L	1.0	-	U	Yes
Trichlorofluoromethane	2.0	ug/L	1.0	-	U	Yes
1,2,4-Trimethylbenzene	2.0	ug/L	1.0	-	U	Yes
Vinyl chloride	1.0	ug/L	1.0	-	U	Yes
m,p-Xylene	1.0	ug/L	1.0	-	U	Yes
o-Xylene	1.0	ug/L	1.0	-	U	Yes
Xylene (total)	1.0	ug/L	1.0	-	U	Yes

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable	
Acetone	10	ug/L	1.0	-	U	Yes	
Benzene	0.5	ug/L	1.0	-	U	Yes	
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes	
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes	
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes	
Bromoform	2.0	ug/L	1.0	-	U	Yes	
Bromomethane	2.0	ug/L	1.0	-	U	Yes	
Butanone (MEK)	10	ug/L	1.0	-	U	Yes	
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes	
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes	
Chlorobenzene	1.4	ug/L	1.0	_	_	Yes	
Chloroethane	1.0	ug/L	1.0	_	U	Yes	
Chloroform	1.0	ug/L	1.0	-	U	Yes	
Chloromethane	5.0	ug/L	1.0	-	Ŭ	Yes	
Cyclohexane	2.0	ug/L	1.0	-	Ū	Yes	
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	_	Ü	Yes	
Dibromochloromethane	1.0	ug/L	1.0	_	Ū	Yes	
1,2-Dibromoethane	1.0	ug/L	1.0	_	Ū	Yes	
1,2-Dichlorobenzene	7.3	ug/L	1.0	-	-	Yes	
1,3-Dichlorobenzene	0.30	ug/L	1.0	J	UJ	Yes	
1,4-Dichlorobenzene	1.1	ug/L	1.0	-	-	Yes	
Dichlorodifluoromethane	2.0	ug/L	1.0	_	U	Yes	
1,1-Dichloroethane	1.0	ug/L	1.0	_	Ü	Yes	
1,2-Dichloroethane	1.0	ug/L	1.0	-	Ü	Yes	
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes	
cis-1,2-Dichloroethene	1.4	ug/L	1.0	_	-	Yes	

trans-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes	
1,2-Dichloropropane	1.0	ug/L	1.0	-	U	Yes	
cis-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes	
trans-1,3-Dichloropropene	1.0	ug/L	1.0	7+3	U	Yes	
Ethylbenzene	1.7	ug/L	1.0	-	-	Yes	
Freon 113	1.0	ug/L	1.0		U	Yes	
2-Hexanone	5.0	ug/L	1.0	-	U	Yes	
Isopropylbenzene	52.1	ug/L	1.0	-	-	Yes	
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes	
Methylcyclohexane	5.0	ug/L	1.0	-	U	Yes	
Methyl Tert Butyl Ether	0.56	ug/L	1.0	J	UJ	Yes	
4-Methyl-2-pentanone (MIBK)	2.0	ug/L	1.0	1-1	U	Yes	
Methylene chloride	2.0	ug/L	1.0	-	U	Yes	
Styrene	1.0	ug/L	1.0	-	U	Yes	
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0	(*)	U	Yes	
Tetrachloroethene	1.0	ug/L	1.0	-	U	Yes	
Tetrahydrofuran	10	ug/L	1.0	-	U	Yes	
Toluene	1.0	ug/L	1.0	-	U	Yes	
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes	
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	4	U	Yes	
1,1,1-Trichloroethane	1.0	ug/L	1.0		U	Yes	
1,1,2-Trichloroethane	1.0	ug/L	1.0	-	U	Yes	
Trichloroethene	0.66	ug/L	1.0	J	UJ	Yes	
Trichlorofluoromethane	2.0	ug/L	1.0	7.1	U	Yes	
1,2,4-Trimethylbenzene	2.0	ug/L	1.0	-	Ų	Yes	
Vinyl chloride	1.0	ug/L	1.0	-	U	Yes	
m,p-Xylene	5.8	ug/L	1.0	-	-	Yes	
o-Xylene	0.50	ug/L	1.0	J	UJ	Yes	
Xylene (total)	6.3	ug/L	1.0		-	Yes	

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	0.32	ug/L	1.0	J	UJ	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	0.36	ug/L	1.0	j	UJ	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	-	U	Yes
Cyclohexane	4.0	ug/L	1.0	J	UJ	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	0.27	ug/L	1.0	J	UJ	Yes
1,3-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,4-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes

trans-1,2-Dichloroethene	1.0	ug/L	1.0		U	Vos
1,2-Dichloropropane	1.0	-	1.0	-		Yes
•		ug/L			Ų	Yes
cis-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes
trans-1,3-Dichloropropene	1.0	ug/L	1.0		U	Yes
Ethylbenzene	0.45	ug/L	1.0	J	UJ	Yes
Freon 113	1.0	ug/L	1.0	1-1	U	Yes
2-Hexanone	5.0	ug/L	1.0		U	Yes
Isopropylbenzene	18.6	ug/L	1.0	/_	-	Yes
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes
Methylcyclohexane	4.9	ug/L	1.0	J	UJ	Yes
Methyl Tert Butyl Ether	1.0	ug/L	1.0	-	U	Yes
4-Methyl-2-pentanone(MIBK)	5.0	ug/L	1.0	-	U	Yes
Methylene chloride	2.0	ug/L	1.0	-	U	Yes
Styrene	1.0	ug/L	1.0	(4)	U	Yes
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0		U	Yes
Tetrachloroethene	1.0	ug/L	1.0	121	U	Yes
Tetrahydrofuran	10	ug/L	1.0	-	Ų	Yes
Toluene	0.42	ug/L	1.0	J	UJ	Yes
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,1,1-Trichloroethane	1.0	ug/L	1.0	17.	U	Yes
1,1,2-Trichloroethane	1.0	ug/L	1.0	3-1	U	Yes
Trichloroethene	1.0	ug/L	1.0	-	U	Yes
Trichlorofluoromethane	2.0	ug/L	1.0	-	U	Yes
1,2,4-Trimethylbenzene	0.83	ug/L	1.0	J	UJ	Yes
Vinyl chłoride	1.0	ug/L	1.0	-	U	Yes
m,p-Xylene	0.61	ug/L	1.0	J	UJ	Yes
o-Xylene	0.98	ug/L	1.0	J	UJ	Yes
Xylene (total)	1.6	ug/L	1.0	-	-	Yes

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	7.4	ug/L	1.0	-	-	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	0.36	ug/L	1.0	J	UJ	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	-	U	Yes
Cyclohexane	5.0	ug/L	1.0	-	U	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	0.48	ug/L	1.0	J	UJ	Yes
1,3-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,4-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes

cis-1,2-Dichloroethene	0.29	ug/L	1.0	J	UJ	Yes
trans-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloropropane	1.0	ug/L	1.0	-	U	Yes
cis-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes
trans-1,3-Dichloropropene	1.0	ug/L	1.0	72	U	Yes
Ethylbenzene	1040	ug/L	20	-	-	Yes
Freon 113	1.0	ug/L	1.0		U	Yes
2-Hexanone	5.0	ug/L	1.0	_	U	Yes
Isopropylbenzene	20.8	ug/L	1.0	-	-	Yes
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes
Methylcyclohexane	5.0	ug/L	1.0		U	Yes
Methyl Tert Butyl Ether	11.8	ug/L	1.0	-	-	Yes
4-Methyl-2-pentanone(MIBK)	5.0	ug/L	1.0	-	U	Yes
Methylene chloride	2.0	ug/L	1.0		U	Yes
Styrene	1.0	ug/L	1.0	-	U	Yes
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0	-	U	Yes
Tetrachloroethene	1.0	ug/L	1.0	-	U	Yes
Tetrahydrofuran	11.2	ug/L	1.0		U	Yes
Toluene	0.25	ug/L	1.0	J	UJ	Yes
1,2,3-Trichlorobenzene	1.0	ug/L	1.0		U	Yes
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,1,1-Trichloroethane	1.0	ug/L	1.0	(4)	U	Yes
1,1,2-Trichloroethane	1.0	ug/L	1.0	-	U	Yes
Trichloroethene	1.0	ug/L	1.0	72	U	Yes
Trichlorofluoromethane	2.0	ug/L	1.0	-	U	Yes
1,2,4-Trimethylbenzene	1.0	ug/L	1.0	100	U	Yes
Vinyl chloride	1.0	ug/L	1.0	-	U	Yes
m,p-Xylene	2480	ug/L	1.0	-	-	Yes
o-Xylene	0.42	ug/L	1.0	1	UJ	Yes
Xylene (total)	2480	ug/L	1.0	141	-	Yes

: 3 ·

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	0.50	ug/L	1.0	-	U	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	0.34	ug/L	1.0	J	UJ	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	-	U	Yes
Cyclohexane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	12.2	ug/L	1.0	-	-	Yes
1,3-Dichlorobenzene	0.49	ug/L	1.0	J	UJ	Yes
1,4-Dichlorobenzene	2.5	ug/L	1.0	-	-	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes

1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	2	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	2	U	Yes
58.5	ug/L	1.0		-	Yes
5.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	2	U	Yes
5.0	ug/L	1.0	-	U	Yes
5.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	*	U	Yes
5.0	ug/L	1.0	7.5	U	Yes
2.0	ug/L	1.0	27	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	7	U	Yes
1.0	ug/L	1.0	25	U	Yes
10	ug/L	1.0	5.00	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	02	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0		U	Yes
2.0	ug/L	1.0		U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	17	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	-	U	Yes
1.0	ug/L	1.0	-	U	Yes
	1.0 1.0 1.0 58.5 5.0 1.0 5.0 5.0 2.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1	1.0 ug/L 1.0 ug/L 1.0 ug/L 1.0 ug/L 1.0 ug/L 58.5 ug/L 5.0 ug/L 1.0 ug/L 5.0 ug/L 5.0 ug/L 1.0 ug/L	1.0 ug/L 1.0 1.0 ug/L 1.0 1.0 ug/L 1.0 1.0 ug/L 1.0 58.5 ug/L 1.0 5.0 ug/L 1.0 1.0 ug/L 1.0	1.0 ug/L 1.0 - 58.5 ug/L 1.0 - 5.0 ug/L 1.0 - 5.0 ug/L 1.0 - 5.0 ug/L 1.0 - 5.0 ug/L 1.0 - 1.0 ug/L 1.0 -	1.0 ug/L 1.0 - U 58.5 ug/L 1.0 - U 5.0 ug/L 1.0 - U 1.0 ug/L 1.0 - U

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016

Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	0.50	ug/L	1.0	•	U	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	0.34	ug/L	1.0	J	UJ	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	-	U	Yes
Cyclohexane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	12.7	ug/L	1.0	-	U	Yes
1,3-Dichlorobenzene	0.46	ug/L	1.0	J	UJ	Yes
1,4-Dichlorobenzene	2.5	ug/L	1.0	-	U	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	•	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes

trans-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes	
1,2-Dichloropropane	1.0	ug/L	1.0	2	U	Yes	
cis-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes	
trans-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes	
Ethylbenzene	1.0	ug/L	1.0	2	U	Yes	
Freon 113	44.0	ug/L	1.0	-	_	Yes	
2-Hexanone	5.0	ug/L	1.0		U	Yes	
Isopropylbenzene	1.0	ug/L	1.0	2	U	Yes	
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes	
Methylcyclohexane	1.0	ug/L	1.0	-	U	Yes	
Methyl Tert Butyl Ether	5.0	ug/L	1.0	-	U	Yes	
4-Methyl-2-pentanone(MIBK)	2.0	ug/L	1.0		U	Yes	
Methylene chloride	1.0	ug/L	1.0	-	U	Yes	
Styrene	1.0	ug/L	1.0	-	U	Yes	
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0		U	Yes	
Tetrachloroethene	10	ug/L	1.0	2	U	Yes	
Tetrahydrofuran	1.0	ug/L	1.0		U	Yes	
Toluene	1.0	ug/L	1.0	-	U	Yes	
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes	
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	Ų	Yes	
1,1,1-Trichloroethane	1.0	ug/L	1.0	-	Ų	Yes	
1,1,2-Trichloroethane	1.0	ug/L	1.0	-	U	Yes	
Trichloroethene	2.0	ug/L	1.0	-	U	Yes	
Trichlorofluoromethane	1.0	ug/L	1.0	_	U	Yes	
1,2,4-Trimethylbenzene	1.0	ug/L	1.0		U	Yes	
Vinyl chloride	1.0	ug/L	1.0	-	U	Yes	
m,p-Xylene	1.0	ug/L	1.0		U	Yes	
o-Xylene	1.0	ug/L	1.0		U	Yes	
Xylene (total)	1.0	ug/L	1.0	-	U	Yes	

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units D	Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	-	Yes
Benzene	0.50	ug/L	1.0	-	-	Yes
Benzyi Chloride	5.0	ug/L	1.0	-	-	Yes
Bromochloromethane	1.0	ug/L	1.0	-	-	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	-	Yes
Bromoform	2.0	ug/L	1.0	-	-	Yes
Bromomethane	2.0	ug/L	1.0	-	-	Yes
Butanone (MEK)	10	ug/L	1.0	-	-	Yes
Carbon disulfide	2.0	ug/L	1.0	-	-	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	-	Yes
Chlorobenzene	0.34	ug/L	1.0	-	-	Yes
Chloroethane	1.0	ug/L	1.0	-	-	Yes
Chloroform	1.0	ug/L	1.0	-	-	Yes
Chloromethane	5.0	ug/L	1.0	-	-	Yes
Cyclohexane	1.0	ug/L	1.0	-	-	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	-	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	-	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	-	Yes
1,2-Dichlorobenzene	1.0	ug/L	1.0	-	-	Yes
1,3-Dichlorobenzene	1.0	ug/L	1.0	-	-	Yes
1,4-Dichlorobenzene	1.0	ug/L	1.0	-	-	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	-	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	-	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	-	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	-	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	-	Yes

trans-1,2-Dichloroethene	1.0	ug/L	1.0		-	Yes
1,2-Dichloropropane	1.0	ug/L	1.0	29	25	Yes
cis-1,3-Dichloropropene	1.0	ug/L	1.0	-	2.4	Yes
trans-1,3-Dichloropropene	1.0	ug/L	1.0	-	-	Yes
Ethylbenzene	1.0	ug/L	1.0	23	-	Yes
Freon 113	5.0	ug/L	1.0	5.5	-	Yes
2-Hexanone	5.0	ug/L	1.0	-	-	Yes
Isopropylbenzene	1.0	ug/L	1.0	-2	-	Yes
Methyl Acetate	5.0	ug/L	1.0	77	-	Yes
Methylcyclohexane	1.0	ug/L	1.0	2.5	(12)	Yes
Methyl Tert Butyl Ether	5.0	ug/L	1.0	*	-	Yes
4-Methyl-2-pentanone(MIBK)	2.0	ug/L	1.0	71	-	Yes
Methylene chloride	1.0	ug/L	1.0			Yes
Styrene	1.0	ug/L	1.0		-	Yes
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0	-	-	Yes
Tetrachloroethene	10	ug/L	1.0	-		Yes
Tetrahydrofuran	1.0	ug/L	1.0	-	-	Yes
Toluene	1.0	ug/L	1.0	-	-	Yes
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	-	-	Yes
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	-	Yes
1,1,1-Trichloroethane	1.0	ug/L	1.0	2	-	Yes
1,1,2-Trichloroethane	1.0	ug/L	1.0	-	-	Yes
Trichloroethene	2.0	ug/L	1.0	-		Yes
Trichlorofluoromethane	1.0	ug/L	1.0	-		Yes
1,2,4-Trimethylbenzene	1.0	ug/L	1.0			Yes
Vinyl chloride	1.0	ug/L	1.0	-	-	Yes
m,p-Xylene	1.0	ug/L	1.0	-	-	Yes
o-Xylene	1.0	ug/L	1.0	2.5	(170)	Yes
Xylene (total)	1.0	ug/L	1.0	-	-	Yes

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	0.50	ug/L	1.0	-	U	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	1.0	ug/L	1.0	-	U	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	-	U	Yes
Cyclohexane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,3-Dichlorobenzene	1.0	ug/L	1.0	-	Ų	Yes
1,4-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes

trans-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes	
1,2-Dichloropropane	1.0	ug/L	1.0	-	U	Yes	
cis-1,3-Dichloropropene	1.0	ug/L	1.0	2	U	Yes	
trans-1,3-Dichloropropene	1.0	ug/L	1.0	5	U	Yes	
Ethylbenzene	1.0	ug/L	1.0	-	U	Yes	
Freon 113	1.0	ug/L	1.0	-	U	Yes	
2-Hexanone	5.0	ug/L	1.0		U	Yes	
Isopropylbenzene	1.0	ug/L	1.0	2	U	Yes	
p-Isopropyltoluene	2.0	ug/L	1.0	-	U	Yes	
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes	
Methylcyclohexane	5.0	ug/L	1.0	2	U	Yes	
Methyl Tert Butyl Ether	2.1	ug/L	1.0		-	Yes	
4-Methyl-2-pentanone(MIBK)	5.0	ug/L	1.0	-	U	Yes	
Methylene chloride	2.0	ug/L	1.0	-	U	Yes	
Styrene	1.0	ug/L	1.0	-	U	Yes	
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0	2	U	Yes	
Tetrachloroethene	1.0	ug/L	1.0	-	U	Yes	
Tetrahydrofuran	10	ug/L	1.0	-	U	Yes	
Toluene	1.0	ug/L	1.0	2	U	Yes	
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes	
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes	
1,1,1-Trichloroethane	1.0	ug/L	1.0		U	Yes	
1,1,2-Trichloroethane	1.0	ug/L	1.0	-	U	Yes	
Trichloroethene	1.0	ug/L	1.0	-	U	Yes	
Trichlorofluoromethane	2.0	ug/L	1.0	*	U	Yes	
1,2,4-Trimethylbenzene	1.0	ug/L	1.0	7.	U	Yes	
Vinyl chloride	1.0	ug/L	1.0	-	U	Yes	
m,p-Xylene	1.0	ug/L	1.0	*.	U	Yes	
o-Xylene	1.0	ug/L	1.0	+	U	Yes	
Xylene (total)	1.0	ug/L	1.0	2	U	Yes	

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

Analyte Name	Result	Units (Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	0.50	ug/L	1.0	-	U	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	0.30	ug/L	1.0	J	UJ	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	-	U	Yes
Cyclohexane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,3-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,4-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes

trans-1,2-Dichloroethene	1.0	ug/L	1.0		U	Yes	
1,2-Dichloropropane	1.0	ug/L	1.0	-	U	Yes	
cis-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes	
trans-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes	
Ethylbenzene	1.0	ug/L	1.0		U	Yes	
Freon 113	1.0	ug/L	1.0	-	U	Yes	
2-Hexanone	5.0	ug/L	1.0	-	U	Yes	
Isopropylbenzene	0.91	ug/L	1.0	J	UJ	Yes	
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes	
Methylcyclohexane	5.0	ug/L	1.0	3.7	U	Yes	
Methyl Tert Butyl Ether	4.5	ug/L	1.0	-	-	Yes	
4-Methyl-2-pentanone(MIBK)	5.0	ug/L	1.0		U	Yes	
Methylene chloride	2.0	ug/L	1.0	0.70	U	Yes	
Styrene	1.0	ug/L	1.0	10-1	U	Yes	
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0	3.5	U	Yes	
Tetrachloroethene	1.0	ug/L	1.0	-	U	Yes	
Tetrahydrofuran	10	ug/L	1.0	-	U	Yes	
Toluene	1.0	ug/L	1.0		U	Yes	
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	7.27	U	Yes	
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes	
1,1,1-Trichloroethane	1.0	ug/L	1.0		U	Yes	
1,1,2-Trichloroethane	1.0	ug/L	1.0	-	U	Yes	
Trichloroethene	1.0	ug/L	1.0	-	U	Yes	
Trichlorofluoromethane	2.0	ug/L	1.0	-	U	Yes	
1,2,4-Trimethylbenzene	1.0	ug/L	1.0	-	U	Yes	
Vinyl chloride	1.0	ug/L	1.0	-	U	Yes	
m,p-Xylene	1.0	ug/L	1.0	-	U	Yes	
o-Xylene	1.0	ug/L	1.0	-	U	Yes	
Xylene (total)	1.0	ug/L	1.0	-	U	Yes	

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
Acetone	10	ug/L	1.0	-	U	Yes
Benzene	0.50	ug/L	1.0	-	U	Yes
Benzyl Chloride	5.0	ug/L	1.0	-	U	Yes
Bromochloromethane	1.0	ug/L	1.0	-	U	Yes
Bromodichloromethane	1.0	ug/L	1.0	-	U	Yes
Bromoform	2.0	ug/L	1.0	-	U	Yes
Bromomethane	2.0	ug/L	1.0	-	U	Yes
Butanone (MEK)	10	ug/L	1.0	-	U	Yes
Carbon disulfide	2.0	ug/L	1.0	-	U	Yes
Carbon tetrachloride	1.0	ug/L	1.0	-	U	Yes
Chlorobenzene	1.0	ug/L	1.0	-	U	Yes
Chloroethane	1.0	ug/L	1.0	-	U	Yes
Chloroform	1.0	ug/L	1.0	-	U	Yes
Chloromethane	5.0	ug/L	1.0	~	U	Yes
Cyclohexane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromo-3-chloropropane	1.0	ug/L	1.0	-	U	Yes
Dibromochloromethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dibromoethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,3-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,4-Dichlorobenzene	1.0	ug/L	1.0	-	U	Yes
Dichlorodifluoromethane	2.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,2-Dichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1-Dichloroethene	1.0	ug/L	1.0	-	U	Yes
cis-1,2-Dichloroethene	1.0	ug/L	1.0	-	U	Yes

trans-1,2-Dichloroethene	1.0	ug/L	1.0	~	U	Yes
1,2-Dichloropropane	1.0	ug/L	1.0	-	U	Yes
cis-1,3-Dichloropropene	1.0	ug/L	1.0	21	U	Yes
trans-1,3-Dichloropropene	1.0	ug/L	1.0	-	U	Yes
Ethylbenzene	1.0	ug/L	1.0	-	U	Yes
Freon 113	1.0	ug/L	1.0	-	U	Yes
2-Hexanone	5.0	ug/L	1.0		U	Yes
Isopropylbenzene	1.0	ug/L	1.0	4	Ų	Yes
p-Isopropyltoluene	2.0	ug/L	1.0		U	Yes
Methyl Acetate	5.0	ug/L	1.0	-	U	Yes
Methylcyclohexane	5.0	ug/L	1.0		U	Yes
Methyi Tert Butyl Ether	1.9	ug/L	1.0		-	Yes
4-Methyl-2-pentanone(MIBK)	5.0	ug/L	1.0	-	U	Yes
Methylene chloride	2.0	ug/L	1.0	-	U	Yes
Styrene	1.0	ug/L	1.0	-	U	Yes
1,1,2,2-Tetrachloroethane	1.0	ug/L	1.0	-	U	Yes
Tetrachloroethene	1.0	ug/L	1.0	-	U	Yes
Tetrahydrofuran	10	ug/L	1.0	-	U	Yes
Toluene	1.0	ug/L	1.0	-	U	Yes
1,2,3-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,2,4-Trichlorobenzene	1.0	ug/L	1.0	-	U	Yes
1,1,1-Trichloroethane	1.0	ug/L	1.0	-	U	Yes
1,1,2-Trichloroethane	1.0	ug/L	1.0		U	Yes
Trichloroethene	1.0	ug/L	1.0	-	U	Yes
Trichlorofluoromethane	2.0	ug/L	1.0	-	U	Yes
1,2,4-Trimethylbenzene	1.0	ug/L	1.0		U	Yes
Vinyl chloride	1.0	ug/L	1.0	22	U	Yes
m,p-Xylene	1.0	ug/L	1.0	-	U	Yes
o-Xylene	1.0	ug/L	1.0	-	U	Yes
Xylene (total)	1.0	ug/L	1.0	-	U	Yes

Project Number:_JC16038	_
Date:March_8-9,_2016	_
Shipping date:March_9-10,_2016	
EPA Region:2_	_

REVIEW OF VOLATILE ORGANIC PACKAGE Low/Medium Volatile Data Validation

The following guidelines for evaluating volatile organics were created to delineate required validation actions. This document will assist the reviewer in using professional judgment to make more informed decision and in better serving the needs of the data users. The sample results were assessed according to USEPA data validation guidance documents in the following order of precedence: USEPA Hazardous Waste Support Section SOP No. HW-33A Revision 0 SOM02.2. Low/Medium Volatile Data Validation. July, 2015. The QC criteria and data validation actions listed on the data review worksheets are from the primary guidance document, unless otherwise noted.

otherwise noted.	
The hardcopied (laboratory name)Accutestbeen reviewed and the quality control and performance VOCs included:	data package received has data summarized. The data review for
Lab. Project/SDG No.:JC16038 No. of Samples:10	Sample matrix:Groundwater
Trip blank No.:JC16038-7	The second of th
X Data CompletenessX Holding TimesX GC/MS TuningX Internal Standard PerformanceX Blanks	X Laboratory Control SpikesX Field DuplicatesX CalibrationsX Compound IdentificationsX Compound QuantitationX Quantitation Limits
Definition of Qualifiers: J- Estimated results J- Compound not detected R- Rejected data JJ- Estimated nondetect Reviewer: Date: April_15,_2016	

DATA REVIEW WORKSHEETS

DATA COMPLETENESS

MISSING INFORMATION	DATE LAB. CONTACTED	DATE RECEIVED

All criteria were met _	_X
Criteria were not met	
and/or see below	

HOLDING TIMES

The objective of this parameter is to ascertain the validity of the results based on the holding time of the sample from time of collection to the time of analysis.

Complete table for all samples and note the analysis and/or preservation not within criteria

SAMPLE ID	DATE SAMPLED	DATE ANALYZED	ρН	ACTION
Samples analyz	ed within method recor	nmended holding time.	Sample	preservation within required
criteria.				procerration manning
Cilicia.				
unteria.				
Cilleria.				
CITERIA.				
Uniteria.				

Criteria

Aqueous samples – 14 days from sample collection for preserved samples (pH \leq 2, 4 \pm 2°C), no air bubbles.

Aqueous samples – 7 days from sample collection for unpreserved samples, 4°C, no air bubbles.

Soil samples- 14 days from sample collection.

Cooler temperature (Criteria: 4 ± 2 °C): 3.6 °C - OK

Actions

Aqueous samples

- a. If there is no evidence that the samples were properly preserved (pH < 2, $T = 4^{\circ}C \pm 2^{\circ}C$), but the samples were analyzed within the technical holding time [7 days from sample collection], no qualification of the data is necessary.
- b. If there is no evidence that the samples were properly preserved, and the samples were analyzed outside of the technical holding time [7 days from sample collection], qualify detects for all volatile compounds as estimated (J) and non-detects as unusable (R).
- c. If the samples were property preserved, and the samples were analyzed within the technical holding time [14 days from sample collection], no qualification of the data is necessary.
- d. If the samples were properly preserved, but were analyzed outside of the technical holding time [14 days from sample collection], qualify detects as estimated (J) and non-detects as unusable (R).
- e. If air bubbles were present in the sample vial used for analysis, qualify detected compounds as estimated (UJ) and non-detected compounds as estimated (UJ).

Non-aqueous samples

- a. If there is no evidence that the samples were properly preserved (T < -7°C or T = 4°C \pm 2°C and preserved with NaHSO₄), but the samples were analyzed within the technical holding time [14 days from sample collection], qualify detects for all volatile compounds as estimated (J) and non-detects as (UJ) or unusable (R) using professional judgment.
- b. If the samples were properly preserved, and the samples were analyzed within the technical holding time [14 days from sample collection], no qualification of the data is necessary.
- c. If there is no evidence that the samples were properly preserved, and the samples were analyzed outside of the technical holding time [14 days from sample collection], qualify detects for all volatile compounds as estimated (J) and non-detects as unusable (R).
- d. If the samples were properly preserved, but were analyzed outside of the technical holding time [14 days from sample collection], qualify detects as estimated (J) and non-detects as unusable (R).

Qualify TCLP/SPLP samples

- a. If the TCLP/SPLP ZHE procedure is performed within the extraction technical holding time of 14 days, detects and non-detects should not be qualified.
- b. If the TCLP/SPLP ZHE procedure is performed outside the extraction technical holding time of 14 days, qualify detects as estimated (J) and non-detects as unusable (R).
- c. If TCLP/SPLP aqueous samples and TCLP/SPLP leachate samples are analyzed within the technical holding time of 7 days, detects and non-detects should not be qualified.
- d. If TCLP/SPLP aqueous samples and TCLP/SPLP leachate samples are analyzed outside of the technical holding time of 7 days, qualify detects as estimated (J) and non-detects as unusable (R).

Table 1. Holding Time Actions for Low/Medium Volatile Analyses - Summary

			Action		
Matrix	Preserved	Criteria Detected Associate Compoun		Non-Detected Associated Compounds	
	No	≤ 7 days	No qualification		
.	No	> 7 days	J	R	
Aqueous	Yes	≤ 14 days	No qualification		
	Yes	> 14 days	J	R	
	No	≤ 14 days	J	Professional judgment, UJ or R	
Non-Aqueous	Yes	≤ 14 days	Noq	ualification	
	Yes/No	> 14 days	J	R	
TCLP/SPLP	Yes	≤ 14 days	No qualification		
TCLP/SPLP	No	> 14 days	J	R	

TCLP/SPLP	ZHE performed within the 14-day technical holding time	No qu	alification
TCLP/SPLP	ZHE performed outside the 14-day technical holding time	J R	
TCLP/SPLP aqueous & TCLP/SPLP leachate	Analyzed within 7 days	No qualification	
TCLP/SPLP aqueous & TCLP/SPLP leachate	Analyzed outside 7 days	J	R
Sample temperature outside 4°C ± 2°C upon receipt at the laboratory		Use professional judgment	
Holding times g	rossly exceeded	J R	

All criteria were met	_
Criteria were not met see below _	20

GC/MS TUNING

The assessment of the tuning results is to determine if the sample instrumentation is within the standard tuning QC limits

__X___The BFB performance results were reviewed and found to be within the specified criteria.
__X___BFB tuning was performed for every 12 hours of sample analysis.

NOTES: All mass spectrometer instrument conditions must be identical to those used during the sample analysis. Background subtraction actions resulting in spectral distortions for the sole purpose of meeting the method specifications are contrary to the Quality Assurance (QA) objectives, and are therefore unacceptable.

NOTES: No data should be qualified based on BFB failure. Instances of this should be noted in the narrative.

All ion abundance ratios must be normalized to m/z 95, the nominal base peak, even though the ion abundance of m/z 174 may be up to 120% that of m/z 95.

Actions:

If samples are analyzed without a preceding valid instrument performance check, qualify all data in those samples as unusable (R).

If ion abundance criteria are not met, professional judgment may be applied to determine to what extent the data may be utilized. When applying professional judgment to this topic, the most important factors to consider are the empirical results that are relatively insensitive to location on the chromatographic profile and the type of instrumentation. Therefore, the critical ion abundance criteria for BFB are the m/z 95/96, 174/175, 174/176, and 176/177 ratios. The relative abundances of m/z 50 and 75 are of lower importance. This issue is more critical for Tentatively Identified Compounds (TICs) than for target analytes.

Note: State in the Data Review Narrative, decisions to use analytical data associated with BFB instrument performance checks not meeting contract requirements.

Note: Verify that that instrument instrument performance check criteria were achieved using techniques described in Low/Medium Volatiles Organic Analysis, Section II.D.5 of the SOM02.2 NFG, obtain additional information on the instrument performance checks. Make sure that background subtraction was performed from the BFB peak and not from background subtracting from the solvent front or from another region of the chromatogram.

DATA REVIEW WORKSHEETS

Use professional judgment to determine whether associated data should be qualified based on the spectrum of the mass calibration compound.			
List	the	samples	affected:
If mass calibrati	on is in error, all associated o	lata are rejected.	

DATA REVIEW WORKSHEETS

All criteria were met _X
Criteria were not met
and/or see below

CALIBRATION VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

Dat	Dates of continuing (initial) calibration:_03/04/16				
Dat				02/26/16	
Dates of continuing calibration: 03/17/16 03/17/16		17/1603/	03/15/16: 03/16/15		
Inst	rument ID num	bers: GCMSA		GCMSU	
			Aqı		
DATE	LAB FILE	CRITERIA OUT	COMPOUND	SAMPLES	
	ID#	RFs, %RSD, %D, r		AFFECTED	
			ithin the required criteria.		
		•	outside method performa		
_		•	ck verification not include	d in data package. No	
action taken	, professional j	udgment.			
	<u> </u>				

Criteria

The analyte calibration criteria in the following Table must be obtained. Analytes not meeting the criteria are qualified.

A separate worksheet should be filled for each initial curve

Initial Calibration - Table 2. RRF, %RSD, and %D Acceptance Criteria for Initial Calibration and CCV for Low/Medium Volatile Analysis

Analyte	Minimum RRF	Maximum %RSD	Opening Maximum %D ¹	Closing Maximum %D
Dichlorodifluoromethane	0.010	25.0	=40.0	±50.0
Chloromethane	0.010	20.0	=30.0	=50.0
Vinyl chloride	0.010	20.0	=25.0	±50.0
Bromomethane	0.010	40.0	=30.0	±50.0
Chloroethane	0.010	40.0	±25.0	±50.0
Trichlorofluoromethane	0.010	40.0	±30.0	±50.0
1.1-Dichloroethene	0.060	20.0	±20.0	=25.0
1,1,2-Trichloro-1,2,2-trifluoroethane	0.050	25.0	±25.0	±50.0
Acetone	0.010	40.0	±40.0	±50.0
Carbon disulfide	0.100	20.0	±25.0	±25.0
Methyl acetate	0.010	40.0	±40.0	±50.0
Methylene chloride	0.010	40.0	±30.0	±50.0
trans-1.2-Dichloroethene	0.100	20.0	=20.0	±25.0
Methyl tert-butyl ether	0.100	40.0	±25.0	±50.0
1.1-Dichloroethane	0.300	20.0	=20.0	±25.0
cis-1,2-Dichloroethene	0.200	20.0	=20.0	=25.0
2-Butanone	0.010	40.0	=40.0	±50.0
Bromochloromethane	0.100	20.0	=20.0	=25.0
Chloroform	0.300	20.0	±20.0	±25.0
1.1.1-Trichloroethane	0.050	20.0	±25.0	±25.0
Cyclohexane	0.010	40.0	±25.0	±50.0
Carbon tetrachloride	0.100	20.0	=25.0	±25.0
Benzene	0.200	20.0	≐20.0	±25.0
1.2-Dichloroethane	0.070	20.0	=20.0	±25.0
Trichloroethene	0.200	20.0	±20.0	±25.0
Methylcyclohexane	0.050	40.0	±25.0	±50.0
1.2-Dichloropropane	0.200	20.0	±20.0	±25.0
Bromodichloromethane	0.300	20.0	±20.0	±25.0
cis-1,3-Dichloropropene	0.300	20.0	±20.0	±25.0
4-Methyl-2-pentanone	0.030	25.0	=30.0	±50.0
Toluene	0.300	20.0	±20.0	±25.0
trans-1,3-Dichloropropene	0.200	20.0	=20.0	±25.0
1.1.2-Trichloroethane	0,200	20.0	=20.0	±25.0
Tetrachloroethene	0.100	20.0	±20,0	±25.0
2-Hexanone	0.010	40.0	=40.0	±50.0
Dibromochloromethane	0.200	20.0	±20.0	±25.0
1.2-Dibromoethane	0.200	20.0	=20.0	±25.0
Chlorobenzene	0.400	20.0	±20.0	±25.0
Ethylbenzene	0.400	20.0	=20.0	±25.0

Analyte	Minimum RRF	Maximum %RSD	Opening Maximum %D ¹	Closing Maximum
ni.p-Xylene	0.200	20.0	±20.0	±25.0
o-Xylene	0.200	20.0	±20.0	±25.0
Styrene	0,200	20.0	±20.0	±25.0
Bromoform	0.100	20.0	=25.0	±50.0
Isopropylbenzene	0.400	20.0	±25.0	±25.0
1.1.2.2-Tetrachloroethane	0.200	20.0	±25.0	±25.0
1.3-Dichlorobenzene	0.500	20.0	=20.0	±25.0
1.4-Dichlorobenzene	0.600	20.0	=20.0	±25.0
1.2-Dichlorobenzene	0,600	20.0	=20.0	±25.0
1.2-Dibromo-3-chloropropane	0.010	25.0	±30.0	±50.0
1.2.4-Trichlorobenzene	0.400	20.0	±30.0	±50.0
1.2.3-Trichlorobenzene	0.400	25.0	=30.0	±50.0
Deuterated Monitoring Compound	·			
Vinyl chloride-di	0.010	20.0	=30.0	±50,0
Chloroethane-ds	0.010	40.0	±30.0	±50.0
1.1-Dichloroethene-da	0.050	20.0	±25.0	±25,0
2-Butanone-ds	0.010	40.0	=40.0	±50.0
Chloroform-d	0,300	20.0	±20.0	±25.0
1.2-Dichloroethane-da	0.060	20.0	±25.0	±25.0
Benzene-de	0.300	20.0	±20.0	±25,0
1.2-Dichloropropane-da	0,200	20.0	≐20.0	±25.0
Toluene-ds	0.300	20.0	=20.0	±25.0
trans-1.3-Dichloropropene-d4	0.200	20.0	=20.0	±25.0
2-Hexanone-ds	0.010	40.0	=40.0	±50.0
1.1.2.2-Tetrachloroethane-d2	0.200	20.0	±25.0	±25.0
1.2-Dichlorobenzene-d4	0.400	20.0	±20.0	±25.0

If a closing CCV is acting as an opening CCV, all target analytes and DMCs must meet the requirements for an opening CCV.

Actions:

- 1. If any volatile target compound has an RRF value less than the minimum in the table, use professional judgment for detects, based on mass spectral identification, to qualify the data as estimated (J+or R).
 - a. If any volatile target compound has an RRF value less than the minimum criterion, qualify non-detected compounds as unusable (R).
 - b. If any of the volatile target compounds listed in the Table has %RSD greater than the criteria, qualify detects as estimated (J), and non-detected compounds using professional judgment.
 - c. If the volatile target compounds meet the acceptance criteria for RRF and the %RSD, no qualification of the data is necessary.

- d. No qualification of the data is necessary on the DMC RRF and %RSD data alone. Use professional judgment and follow the guidelines in Action 2 to evaluate the DMC RRF and %RSD data in conjunction with the DMC recoveries to determine the need for qualification of data.
- 2. At the reviewer's discretion, and based on the project-specific Data Quality Objectives (DQOs), a more in-depth review may be considered using the following guidelines:
 - a. If any volatile target compound has a %RSD greater than the maximum criterion in the Table, and if eliminating either the high or the low-point of the curve does not restore the %RSD to less than or equal to the required maximum:
 - Qualify detects for that compound(s) as estimated (J).
 - ii. Qualify non-detected volatile target compounds using professional judgment.
 - b. If the high-point of the curve is outside of the linearity criteria (e.g., due to saturation):
 - i. Qualify detects outside of the linear portion of the curve as estimated (J).
 - ii. No qualifiers are required for detects in the linear portion of the curve.
 - iii. No qualifiers are required for volatile target compounds that were not detected.
 - c. If the low-point of the curve is outside of the linearity criteria:
 - i. Qualify low-level detects in the area of non-linearity as estimated (J).
 - ii. No qualifiers are required for detects in the linear portion of the curve.
 - iii. For non-detected volatile compounds, use the lowest point of the linear portion of the curve to determine the new quantitation limit.

Note: If the laboratory has failed to provide adequate calibration information, inform the Region's designated representative to contact the laboratory and request the necessary information. If the information is not available, the reviewer must use professional judgment to assess the data.

State in the Data Review Narrative, if possible, the potential effects on the data due to calibration criteria exceedance.

Note, for the Laboratory COR action, if calibration criteria are grossly exceeded.

Table. Initial Calibration Actions for Low/Medium Volatile Analysis - Summary

Criteria	Action		
Criteria	Detect	Non-detect Use professional judgment R	
Initial Calibration not performed at specified frequency and sequence	Use professional judgment R		
Initial Calibration not performed at the specified concentrations	3	UJ	
RRF = Minimum RRF in Table for target analyte	Use professional judgment J+ or R	R	
RRF = Minimum RRF in Table for integer analyte	No qualification	No qualification	
*«RSD = Maximum *«RSD in Table for target analyte	J	Use professional judgment	
° ∘RSD :: Maximum ° ∘RSD in Table for target analyte	No qualification	No qualification	

All criteria were met _X	
Criteria were not met	
and/or see below	

Continuing Calibration Verification (CCV)

NOTE: Verify that the CCV was run at the required frequency (an opening and closing CCV must be run within 12-hour period) and the CCV was compared to the correct initial calibration. If the mid-point standard from the initial calibration is used as an opening CCV, verify that the result (RRF) of the mid-point standard was compared to the average RRF from the correct initial calibration.

The closing CCV used to bracket the end of a 12-hour analytical sequence may be used as the opening CCV for the new 12-hour analytical sequence, provided that all the technical acceptance criteria are met for an opening CCV (see criteria show before in the Table). If the closing CCV does not meet the technical acceptance criteria for an opening CCV, then a BFB tune followed by an opening CCV is required and the next 12-hour time period begins with the BFB tune.

All DMCs must meet RRF criteria. No qualification of the data is necessary on the DMCs RRF and %RSD/%D data alone. However, use professional judgment to evaluate the DMC and %RSD/%D data in conjunction with the DMC recoveries to determine the need of qualification the data.

Action:

- 1. If a CCV (opening and closing) was not run at the appropriate frequency, qualify data using professional judgment.
- 2. Qualify all volatile target compounds in Table shown before using the following criteria:
 - a. For an opening CCV, if any volatile target compound has an RRF value less than the minimum criterion, use professional judgment for detects, based on mass spectral identification, to qualify the data as estimated (J) and qualify non-detected compounds as unusable (R).
 - b. For a closing CCV, if any volatile target compound has an RRF value less than the criteria, use professional judgment for detects based on mass spectral identification to qualify the data as estimated (J), and qualify non-detected compounds as unusable (R).
 - c. For an opening CCV, if the Percent Difference value for any of the volatile target compounds is outside the limits in calibration criteria Table shown before, qualify detects as estimated (J) and non-detected compounds as estimated (UJ).
 - For a closing CCV, if the Percent Difference value for any volatile target compound is outside the limits in calibration criteria table, qualify detects as estimated (J) and non-detected compounds as estimated (UJ).
 - e. If the volatile target compounds meet the acceptable criteria for RRF and the Percent Difference, no qualification of the data is necessary.

DATA REVIEW WORKSHEETS

f. No qualification of the data is necessary on the DMC RRF and the Percent Difference data alone. Use professional judgment to evaluate the DMC RRF and Percent Difference data in conjunction with the DMC recoveries to determine the need for qualification of data.

Notes: If the laboratory has failed to provide adequate calibration information, inform the Region's designated representative to contact the laboratory and request the necessary information. If the information is not available, the reviewer must use professional judgment to assess the data.

State in the Data Review Narrative, if possible, the potential effects on the data due to calibration criteria exceedance.

Note, for Contract Laboratory COR action, if calibration criteria are grossly exceeded.

Table. Continuing Calibration Actions for Low/Medium Volatile Analysis - Summary

Criteria for Opening	Criteria for	A	ction
CCI.	Closing CCV	Detect	Non-detect
CCV not performed at required frequency	CCV not performed at required frequency	Use professional judgment R	Use professional judgment R
CCV not performed at specified concentration	CCV not performed at specified concentration	Use professional judgment	Use professional judgment
RRF - Minimum RRF in Table 2 for target analyte	RRF < Minimum RRF in Table for target analyte	Use professional judgment J or R	R
RRF :: Minimum RRF in Table 2 for target analyte	RRF ≥ Minimum RRF in Table for target analyte	No qualification	No qualification
%D outside the Opening Maximum %D limits in Table 2 for target analyte	°oD outside the Closing Maximum °oD limits in Table for target analyte	J	UJ
⁹ •D within the inclusive Opening Maximum ⁹ •D limits in Table 2 for target analyte	%D within the inclusive Closing Maximum %D limits in Table for target madyte	No qualification	No qualification

All criteria were metX	
Criteria were not met	
and/or see below	

BLANK ANALYSIS RESULTS (Sections 1 & 2)

The assessment of the blank analysis results is to determine the existence and magnitude of contamination problems. The criteria for evaluation of blanks apply only to blanks associated with the samples, including trip, equipment, and laboratory blanks. If problems with any blanks exist, all data associated with the case must be carefully evaluated to determine whether or not there is an inherent variability in the data for the case, or if the problem is an isolated occurrence not affecting other data.

List the contamination in the blanks below. High and low levels blanks must be treated separately.

The concentration of a target analyte in any blank must not exceed its Contract Required Quantitation Limit (CRQL) (2x CRQLs for Methylene chloride, Acetone, and 2-Butanone). TIC concentration in any blanks must be $\leq 5.0 \,\mu\text{g/L}$ for water (0.0050 mg/L for TCLP leachate) and $\leq 5.0 \,\mu\text{g/kg}$ for soil matrices.

Laboratory blanks

The method blank, like any other sample in the SDG, must meet the technical acceptance criteria for sample analysis.

DATE ANALYZED	LAB ID	LEVEL! MATRIX	COMPOUND	CONCENTRATION UNITS
No_target_analyt	te_detected_in_	method_blanks	3.	

Field/Equipment/Trip blank

If field or trip blanks are present, the data reviewer should evaluate this data in a similar fashion as the method blanks.

DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS
_No_target_ana _this_data_pack	alytes_detected_ kage	_in_the_trip_blank	No_equipment_blank_a	nalyzed_as_part_of
Field_Biank	-			
03/15/16	JC16038-9	_Aqueous/low	Chlorobenzene	0.30_ug/L
			lsopropyl_benzene	0.91_ug/L
			MTBE	4.5ug/L

Note: No action taken, professional judgment.

All criteria were metX
Criteria were not met
and/or see below

BLANK ANALYSIS RESULTS (Section 3)

Blank Actions

Note:

All fields blank results associated with a particular group of samples (may exceed one per case) must be used to qualify data. Trip blanks are used to qualify only those samples with which they were shipped. Blanks may not be qualified because of contamination in another blank. Field blanks and trip blanks must be qualified for system monitoring compounds, instrument performance criteria, and spectral or calibration QC problems.

Samples taken from a drinking water tap do not have associated field blanks.

When applied as described in the Table below, the contaminant concentration in the blank is multiplied by the sample dilution factor.

Table. Blank and TCLP/SPLP LEB Actions for Low/Medium Volatile Analysis

Blank Type	Blank Result	Sample Result	Action for Samples	
Method, Storage, Field, Trip, TCLP/SPLP LEB, Instrument**	Detects	Not detected	No qualification required	
	< CRQL *	≤ CRQL*	Report CRQL value with a U	
		≥ CRQL*	No qualification required	
	> CRQL *	< CRQL*	Report CRQL value with a U	
		≥ CRQL** and ≤	Report blank value for sample	
		blank concentration	concentration with a U	
		≥ CRQL* and >	No qualification required	
		blank concentration	No quantication required	
	= CRQL*	≤CRQL*	Report CRQL value with a U	
	CRQL	> CRQL*	No qualification required	
	Gross	Detects	Report blank value for sample	
	contamination	Detects	concentration with a U	

^{* 2}x the CRQL for methylene chloride, 2-butanone and acetone.

Action Levels (ALs) should be based upon the highest concentration of contaminant determined in any blank. Do not qualify any blank with another blank. The ALs for samples which have been diluted should be corrected for the sample dilution factor and/or % moisture, where applicable. No positive sample results should be reported unless the concentration of the compound in the samples exceeds the ALs:

^{**} Qualifications based on instrument blank results affect only the sample analyzed immediately after the sample that has target compounds that exceed the calibration range or non-target compounds that exceed 100 µg/L.

DATA REVIEW WORKSHEETS

Notes:

High and low level blanks must be treated separately Compounds qualified "U" for blank contamination are still considered "hits" when qualifying for calibration criteria.

CONTAMINATION SOURCE/LEVEL	COMPOUND	CONC/UNITS	AL/UNITS	SQL	AFFECTED SAMPLES
<u>_</u>					
···					
		+			
		_		_	
	<u> </u>	+			
					
-					

All criteria were met _	X
Criteria were not met	
and/or see below	

DEUTERATED MONITORING COMPOUNDS (DMCs)

Laboratory performance of individual samples is established by evaluation of surrogate spike (DMCs) recoveries. All samples are spiked with surrogate compounds prior to sample analysis. The accuracy of the analysis is measured by the surrogate percent recovery. Since the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the validation of data is frequently subjective and demands analytical experience and professional judgment.

Table. Volatile Deuterated Monitoring Compounds (DMCs) and Recovery Limits

DMC	%R for Water Sample	%R for Soil Sample
Vinyl chloride-d3	60-135	30-150
Chloroethane-d5	70-130	30-150
1.1-Dichloroethene-d2	60-125	45-110
2-Butanone-d5	40-130	20-135
Chloroform-d	70-125	40-150
1.2-Dichloroethane-d4	70-125	70-130
Benzene-d6	70-125	20-135
1.2-Dichloropropane-d6	70-120	70-120
Toluene-d8	80-120	30-130
trans-1,3-	60-125	30-135
Dichloropropene-d4		
2-Hexanone-d5	45-130	20-135
1.1.2.2-	65-120	45-120
Tetrachloroethane-d2		
1.2-Dichlorobenzene-d4	80-120	75-120

NOTE: The recovery limits for any of the compounds listed in the above Table may be expanded at any time during the period of performance if the United States Environmental Protection Agency (EPA) determines that the limits are too restrictive.

Action:

Are recoveries for DMCs in volatile samples and blanks must be within the limits specified in the Table above.

Yes? or No?

NOTE: The recovery limits for any of the compounds listed in the Table above may be expanded at any time during the period of performance if USEPA determines that the limits are too restrictive.

List the DMCs that may fail to meet the recovery limits

Sample ID

Date

DMCs

% Recovery

Action

DMCs recoveries within the required limits. Other non-deuterated surrogates added to the samples within laboratory control limits.

Note: Any sample which has more than 3 DMCs outside the limits must be reanalyzed.

Action:

1. For any recovery greater than the upper acceptance limit:

- a. Qualify detected associated volatile target compounds as estimated high (J+).
- b. Do not qualify non-detected associated volatile target compounds.
- 2. For any recovery greater than or equal to 10%, and less than the lower acceptance limit:
 - a. Qualify detected associated volatile target compounds as estimated low (J-).
 - Qualify non-detected associated volatile target compounds as estimated (UJ).
- 3. For any recovery less than 10%:
 - a. Qualify detected associated volatile target compounds as estimated low (J-).
 - b. Qualify non-detected associated volatile target compounds as unusable (R).
- 4. For any recovery within acceptance limits, no qualification of the data is necessary.
- In the special case of a blank analysis having DMCs out of specification, the reviewer must give special consideration to the validity of associated sample data. The basic concern is whether the blank problems represent an isolated problem with the blank alone, or whether there is a fundamental problem with the analytical process. For example, if one or more samples in the batch show acceptable DMC recoveries, the reviewer may choose to consider the blank problem to be an isolated occurrence. However, even if this judgment allows some use of the affected data, note analytical problems for Contract Laboratory COR action.
- If more than three DMCs are outside of the recovery limits for Low/Medium volatiles analysis and the sample was not reanalyzed, note under Contract Problems/Non-Compliance.

Table. Deuterated Monitoring Compound (DMC) Recovery Actions for Low/Medium Volatiles Analyses – Summary

	Action			
Criteria	Detect Associated Compounds	Non-detected Associated Compounds		
° oR < 10° o	J-	R		
10° o ≤ ° oR < Lower Acceptance Limit	J-	UJ		
Lower Acceptance Limit $\leq 6 _0 R \leq U_{pper}$ Acceptance Limit	No qualification No qualification			
% oR → Upper Acceptance Limit	J _†	No qualification		

DATA REVIEW WORKSHEETS

TABLE. VOLATILE DEUTERATED MONITORING COMPOUNDS (DMCs) AND THE ASSOCIATED TARGET COMPOUNDS

Vinyl chloride-d3 (DMC-1)	Chloroethane-ds (DMC-2)	1,1-Dichloroethene-d: (DMC-3)
Vinyl chloride	Dichlorodifluoromethane Chloromethane	trans-1,2-Dichloroethene cis-1,2-Dichloroethene
	Bromomethane	1.1-Dichloroethene
	Chloroethane Carbon disulfide	
2-Butanone-ds (DMC-4)	Chloroform-d (DMC-5)	1,2-Dichloroethane-d+(DMC-6)
Acetone	1.1-Dichloroethane	Trichlorofluoromethane
2-Butanone	Bromochloromethane	1.1.2-Trichloro-1.2,2-trifluoroethane
	Chloroform	Methyl acetate
	Dibromochloromethane	Methylene chloride
	Bromoform	Methyl-tert-butyl ether
		1.1.1-Trichloroethane Carbon tetrachloride
		1.2-Dibromoethane
	ļ	1.2-Dichloroethane
Benzene-da (DMC-7)	1,2-Dichloropropane-de	Toluene-ds (DMC-9)
Benzene-da (BMC-7)	(DMC-8)	Toluene-ds (DMC-9)
Benzene	Cyclohexane	Trichloroethene
	Methylcyclohexane	Toluene
	1.2-Dichloropropane	Tetrachloroethene
	Bromodichloromethane	Ethylbenzene
		o-Xylene
		m.p-Xylene Styrene
		Isopropylbenzene
trans-1,3-Dichloropropene-da	2-Hexanone-ds (DMC-11)	1,1,2,2-Tetrachloroethane-d:
(DMC-10)		(DMC-12)
cis-1,3-Dichloropropene	4-Methyl-2-pentanone	1.1.2.2Tetrachloroethane
trans-1,3-Dichloropropene	2-Hexanone	1.2-Dibromo-3-chloropropane
1.1.2-Trichloroethane		·
1,2-Dichlorobenzene-da		
(DMC-13)		
Chlorobenzene	1	
1.3-Dichlorobenzene		
1.4-Dichlorobenzene		
1.2-Dichlorobenzene		
1.2.4-Trichlorobenzene		
1.2.3-Trichforobenzene		

All criteria were met _X	
Criteria were not met	
and/or see below	

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD)

This data is generated to determine long term precision and accuracy in the analytical method for various matrices. This data alone cannot be used to evaluate the precision and accuracy of individual samples. If any % R in the MS or MSD falls outside the designated range, the reviewer should determine if there are matrix effects, i.e. LCS data are within the QC limits but MS/MSD data are outside QC limit.

NOTES:

Data for MS and MSDs will not be present unless requested by the Region.

Notify the Contract Laboratory COR if a field or trip blank was used for the MS and MSD.

For a Matrix Spike that does not meet criteria, apply the action to only the field sample used to prepare the Matrix Spike sample. If it is clearly stated in the data validation materials that the samples were taken through incremental sampling or some other method guaranteeing the homogeneity of the sample group, then the entire sample group may be qualified.

1. MS/MSD Recoveries and Precision Criteria

The laboratory should use one MS and a duplicate analysis of an unspiked field sample if target analytes are expected in the sample. If target analytes are not expected, MS/MSD should be analyzed.

List the %Rs, RPD of the compounds which do not meet the criteria.

MS OR MSD COMPOUND % R RPD QC LIMITS ACTION _MS/MSD_%_recovery_and_RPD_within_laboratory_control_limits_except_for_the_followings:	Sample ID:_JC16101-1MS Sample ID:_JC16065-1MS Sample ID:_JC16250-14MS/MSD		Matrix	/Level:Groundw /Level:Groundw /Level:Groundw	ater	
_MS/MSDm&_p-xylene30%/34%42139No_action	_MS/MSD_%_re	_			A .	
	_MS/MSD	m&_p-xylene	30%/34%	6	42139	No_action

Note: No action taken, sample concentration high relative to amount spiked.

MS/MSD criteria apply to the unspiked sample. Unspiked sample belongs to from another data package.

* QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper limit.

If QC limits are not available, use limits of 70 – 130 %.

DATA REVIEW WORKSHEETS

Actions:

1. No qualification of the data is necessary on MS and MSD data alone. However, using professional judgment, the validator may use the MS and MSD results in conjunction with other QC criteria and determine the need for some qualification of the data.

QUALITY	%R < LL	%R > UL
Positive results	J	J
Nondetects results	R	Accept

MS/MSD criteria apply only to the unspiked sample, its dilutions, and the associated MS/MSD samples:

If the % R for the affected compounds were < LL (or 70 %), qualify positive results (J) and nondetects (UJ).

If the % R for the affected compounds were > UL (or 130 %), only qualify positive results (J).

If 25 % or more of all MS/MSD %R were < LL (or 70 %) or if two or more MS/MSD %Rs were < 10%, qualify all positive results (J) and reject nondetects (R).

A separate worksheet should be used for each MS/MSD pair.

All criteria were metX
Criteria were not met
and/or see below

LABORATORY CONTROL SAMPLE (LCS) ANALYSIS

This data is generated to determine accuracy of the analytical method for various matrices.

1. LCS Recoveries Criteria

Where LCS spiked with the same analyte at the same concentrations as the MS/MSD? Yes or No. If no make note in data review memo.

List the %R of compounds which do not meet the criteria

	LCS ID	COMPOUND	% R	QC LIMIT
Recoveri	es_(blank_spike	e)_within_laboratory_control	_limits	

- * QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper limit.
- * If QC limits are not available, use limits of 70 130 %.

Actions:

QUALITY	%R < LL	%R > UL
Positive results	J	J
Nondetects results	R	Accept

All analytes in the associated sample results are qualified for the following criteria.

If 25 % of the LCS recoveries were < LL (or 70 %), qualify all positive results (j) and reject nondetects (R).

If two or more LCS were below 10 %, qualify all positive results as (J) and reject nondetects (R).

2. Frequency Criteria:

Where LCS analyzed at the required frequency and for each matrix? <u>Yes</u> or No. If no, the data may be affected. Use professional judgment to determine the severity of the effect and qualify data accordingly. Discuss any actions below and list the samples affected.

All criteria were met _X
Criteria were not met
and/or see below

IX. FIELD/LABORATORY DUPLICATE PRECISION

Sample IDs:	_JC16038-5/-6	Matrix:_Groundwater
-------------	---------------	---------------------

Field/laboratory duplicates samples may be taken and analyzed as an indication of overall precision. These analyses measure both field and lab precision; therefore, the results may have more variability than laboratory duplicates which only laboratory performance. It is also expected that soil duplicate results will have a greater variance than water matrices due to difficulties associated with collecting identical field duplicate samples.

The project QAPP should be reviewed for project-specific information.

NOTE: In the absence of QAPP guidance for validating data from field duplicates, the following action will be taken.

Identify which samples within the data package are field duplicates. Estimate the relative percent difference (RPD) between the values for each compound. Use professional judgment to note large RPDs (> 50%) in the narrative.

COMPOUND	SQL	SAMPLE CONC.	DUPLICATE CONC.	RPD	ACTION
DDD withi	n require	nd critoria < 50 % fo	r target analytes detect	od in som	
TAI D WIGH	Trequit	Turnena, > 50 % ic	i larget allarytes detect	Eu III Saii	pie and dupilcate.

Actions:

Qualify as estimated positive results (J) and nondetects (UJ) for the compound that exceeded the above criteria. For organics, only the sample and duplicate will be qualified.

If an RPD cannot be calculated because one or both of the sample results is not detected, the following actions are suggested based on professional judgment:

If one sample result is not detected and the other is greater than 5x the SQL qualify (J/UJ).

If one sample value is not detected and the other is greater than 5x the SQL and the SQLs for the sample and duplicate are significantly different, use professional judgment to determine if qualification is appropriate.

If one sample value is not detected and the other is less than 5x, use professional judgment to determine if qualification is appropriate.

If both sample and duplicate results are not detected, no action is needed.

All criteria were met __X__ Criteria were not met and/or see below ____

X. INTERNAL STANDARD PERFORMANCE

The assessment of the internal standard (IS) parameter is used to assist the data reviewer in determining the condition of the analytical instrumentation.

DATE SAMPLE ID IS OUT IS AREA ACCEPTABLE ACTION RANGE

Internal standard area counts within the required criteria.

Action:

- 1. If an internal standard area count for a sample or blank is greater than 200.0% of the area for the associated standard (opening CCV or mid-point standard from initial calibration) (see Table below):
 - a. Qualify detects for compounds quantitated using that internal standard as estimated low (J-).
 - b. Do not qualify non-detected associated compounds.
- 2. If an internal standard area count for a sample or blank is less than 20.0% of the area for the associated standard (opening CCV or mid-point standard from initial calibration):
 - a. Qualify detects for compounds quantitated using that internal standard as estimated high (J+).
 - b. Qualify non-detected associated compounds as unusable (R).
- If an internal standard area count for a sample or blank is greater than or equal to 20.0%, and less than or equal to 200% of the area for the associated standard opening CCV or mid-point standard from initial calibration, no qualification of the data is necessary.
- 4. If an internal standard RT varies by more than 30.0 seconds: Examine the chromatographic profile for that sample to determine if any false positives or negatives exist. For shifts of a large magnitude, the reviewer may consider partial or total rejection of the data for that sample fraction. Detects should not need to be qualified as unusable (R) if the mass spectral criteria are met.
- If an internal standard RT varies by less than or equal to 30.0 seconds, no qualification of the data is necessary.

Note: Inform the Contract Laboratory Program Project Officer (CLP PO) if the internal standard performance criteria are grossly exceeded. Note in the Data Review Narrative potential effects on the data resulting from unacceptable internal standard performance.

DATA REVIEW WORKSHEETS

- 6. If required internal standard compounds are not added to a sample or blank, qualify detects and non-detects as unusable (R).
- 7. If the required internal standard compound is not analyzed at the specified concentration in a sample or blank, use professional judgment to qualify detects and non-detects.

Table. Internal Standard Actions for Low/Medium Volatiles Analyses - Summary

	Act	tion
Criteria	Detected Associated Compounds*	Non-detected Associated Compounds*
Area counts > 200% of 12-hour standard (opening CCV or mid-point standard from initial calibration)	J-	No qualification
Area counts < 20% of 12-hour standard (opening CCV or mid-point standard from initial calibration)	J-	R
Area counts ≥ 50% but ≤ 200% of 12-hour standard (opening CCV or mid-point standard from initial calibration)	No qualification	
RT difference > 30.0 seconds between samples and 12-hour standard (opening CCV or mid-point standard from initial calibration)	R **	R
RT difference ≤ 30.0 seconds between samples and 12-hour standard (opening CCV or mid-point standard from initial calibration)	No qual	ification

^{*} For volatile compounds associated to each internal standard, see TABLE - VOLATILE TARGET ANALYTES, DEUTERATED MONITORING COMPOUNDS WITH ASSOCIATED INTERNAL STANDARDS FOR QUANTITATION in SOM02.2, Exhibit D, available at: http://www.epa.gov/superfund/programs/clp/download/som/som22d.pdf

^{**} Detects should not need to be qualified as unusable (R) if the mass spectral criteria are met.

All criteria were met _	X
Criteria were not met	
and/or see below	

TARGET COMPOUND IDENTIFICATION

Criteria:

Is the Relative Retention Times (RRTs) of reported compounds within ±0.06 RRT units of the standard RRT [opening Continuing Calibration Verification (CCV) or mid-point standard from the initial calibration].

Yes? or No?

List compounds not meeting the criteria described above:

Sample ID	Compounds	Actions

Mass spectra of the sample compound and a current laboratory-generated standard [i.e., the mass spectrum from the associated calibration standard (opening CCV or mid-point standard from initial calibration)] must match according to the following criteria:

- a. All ions present in the standard mass spectrum at a relative intensity greater than 10% must be present in the sample spectrum.
- b. The relative intensities of these ions must agree within $\pm 20\%$ between the standard and sample spectra (e.g., for an ion with an abundance of 50% in the standard spectrum, the corresponding sample ion abundance must be between 30-70%).
- c. lons present at greater than 10% in the sample mass spectrum, but not present in the standard spectrum, must be evaluated by a reviewer experienced in mass spectral interpretation.

List compounds not meeting the criteria described above:

Sample ID	Compounds			Actions	
	 		_		

DATA REVIEW WORKSHEETS

Action:

- The application of qualitative criteria for GC/MS analysis of target compounds requires
 professional judgment. It is up to the reviewer's discretion to obtain additional information
 from the laboratory. If it is determined that incorrect identifications were made, qualify all
 such data as unusable (R).
- Use professional judgment to qualify the data if it is determined that cross-contamination has occurred.
- 3. Note in the Data Review Narrative any changes made to the reported compounds or concerns regarding target compound identifications. Note, for Contract Laboratory COR action, the necessity for numerous or significant changes.

TENTATIVELY IDENTIFIED COMPOUNDS (TICS)

NOTE: Tentatively identified compounds should only be evaluated when requested by a party from outside of the Hazardous Waste Support Section (HWSS).

1	ict	TI	Ce

Sample ID	Compound	Sample ID	Compound

Action:

- 1. Qualify all TIC results for which there is presumptive evidence of a match (e.g. greater than or equal to 85% match) as tentatively identified (NJ), with approximated concentrations. TICs labeled "unknown" are qualified as estimated (J).
- 2. General actions related to the review of TIC results are as follows:
 - a. If it is determined that a tentative identification of a non-target compound is unacceptable, change the tentative identification to "unknown" or another appropriate identification, and qualify the result as estimated (J).
 - b. If all contractually-required peaks were not library searched and quantitated, the Region's designated representative may request these data from the laboratory.
- 3. In deciding whether a library search result for a TIC represents a reasonable identification, use professional judgment. If there is more than one possible match, report the result as "either compound X or compound Y". If there is a lack of isomer specificity, change the TIC result to a nonspecific isomer result (e.g., 1,3,5-trimethyl benzene to trimethyl benzene

DATA REVIEW WORKSHEETS

- isomer) or to a compound class (e.g., 2-methyl, 3-ethyl benzene to a substituted aromatic compound).
- 4. The reviewer may elect to report all similar compounds as a total (e.g., all alkanes may be summarized and reported as total hydrocarbons).
- 5. Target compounds from other fractions and suspected laboratory contaminants should be marked as "non-reportable".
- 6. Other Case factors may influence TIC judgments. If a sample TIC match is poor, but other samples have a TIC with a valid library match, similar RRT, and the same ions, infer identification information from the other sample TIC results.
- 7. Note in the Data Review Narrative any changes made to the reported data or any concerns regarding TIC identifications.
- 8. Note, for Contract Laboratory COR action, failure to properly evaluate and report TICs

All criteria were met	X_
Criteria were not met	
and/or see below	

SAMPLE QUANTITATION AND REPORTED CONTRACT REQUIRED QUANTITATION LIMITS (CRQLS)

Action:

- 1. If any discrepancies are found, the Region's designated representative may contact the laboratory to obtain additional information that could resolve any differences. If a discrepancy remains unresolved, the reviewer must use professional judgment to decide which value is the most accurate. Under these circumstances, the reviewer may determine that qualification of data is warranted. Note in the Data Review Narrative a description of the reasons for data qualification and the qualification that is applied to the data.
- 2. For non-aqueous samples, in the percent moisture is less than 70.0%, no qualification of the data is necessary. If the percent moisture is greater than or equal to 70.0% and less than 90.0%, qualify detects as estimated (J) and non-detects as approximated (UJ). If the percent moisture is greater than or equal to 90.0%, qualify detects as estimated (J) and non-detects as unusable (R) (see Table below).
- 3. Note, for Contract Laboratory COR action, numerous or significant failures to accurately quantify the target compounds or to properly evaluate and adjust CRQLs.
- 4. Results between MDL and CRQL should be qualified as estimated "J".
- 5. Results < MDL should be reported at the CRQL and qualified "U". MDLs themselves are not reported.

Table. Percent Moisture Actions for Low/Medium Volatiles Analysis for Non-Aqueous Samples

Criteria	Action		
	Detected Associated Compounds	Non-detected Associated Compounds	
% Moisture < 70.0	No	qualification	
70.0 < % Moisture < 90.0	J	UJ	
% Moisture > 90.0	J	R	

The sample quantitation evaluation is to verify laboratory quantitation results. In the space below, please show a minimum of one sample calculation:

Sample ID

JC16038-1

Freon 113

RF = 0.279

[] = (38214)(50)/(0.279)(227356) = 30.12 ppb Ok

DATA REVIEW WORKSHEETS

В.	Percent Solids
	List samples which have ≥ 70 % solids

All criteria were met _X_	_
Criteria were not met	
and/or see below	

QUANTITATION LIMITS

A. Dilution performed

20 X	Ethylbenzene and the xylenes outside
	calibration range.
_	

Assessment (DQA).

All criteria were met_	Χ
Criteria were not met	
and/or see below	_

OTHER ISSUES

A.	System Perform	ance	
List sa	mples qualified ba	sed on the degradation of system	performance during simple analysis:
Sampl		Comments	Actions
		Ann northware about d	
Action:			
degrad	led during sample		etermined that system performance has aboratory Program COR any action as a antly affected the data.
В.	Overall Assessme	ent of Data	
List sa	mples qualified ba	sed on other issues:	
	e ID	Comments	Actions
_can_t		sion_purposes	on_of_the_dataResults_are_valid_and_
Action: 1. 2.	qualified based of Write a brief nam Inform the Contra Delivery Group (n the Quality Control (QC) criteria prative to give the user an indication of Laboratory COR the action, any SDG) Narrative. If sufficient inform	any need to qualify data which were not previously discussed. In of the analytical limitations of the data, inconsistency of the data with the Sample mation on the intended use and required include their assessment of the usability of

the data within the given context. This may be used as part of a formal Data Quality

EXECUTIVE NARRATIVE

SDG No:

JC16038

Laboratory:

Accutest, New Jersey

Analysis:

SW846-8270D

Number of Samples:

9

Location:

BMSMC, Former Tank Farm Area

Humacao, PR

SUMMARY:

Eight (8) groundwater samples and one (1) field blank were analyzed for the ABN TCL list following method SW846-8270D; Naphthalene and 1,4-Dioxane were also analyzed by SW846-8270D using the selective ion monitoring (SIM) technique. The sample results were assessed according to USEPA data validation guidance documents in the following order of precedence: EPA Hazardous Waste Support Section, SOP HW-35A, July 2015 – Revision 0. Semivolatile Data Validation. The QC criteria and data validation actions listed on the data review worksheets are from the primary guidance document, unless otherwise noted.

Results are valid and can be used for decision making purposes.

Critical issues:

None

Major:

None

Minor:

- 1. Closing calibration verification not included in date package. None of the results were qualified, professional judgment.
- 2. Several analytes did not meet the method required criteria for % difference in the continuing calibration verification, but were within the validation guidance document %D required criteria. No action taken.
- 3. bis(2-ethylhexyl)phthalate detected in the field blank at a concentration of 1.2 ug/L. No action taken, professional judgment.
- 4. % recovery for 2-Fluorobiphenyl (surrogate) outside laboratory control limits due to dilution. No action taken, professional judgment.
- 5. 1,4-Dioxane MS/MSD % recovery outside the laboratory control limits. No action taken, high level of sample relative to amount spiked.
- 6. Results for 2-methylnaphthalene and naphthalene qualified as estimated (J) in samples JC16038-5 and JC16038-6, RPD outside the required criteria (< 50 %) in the field duplicate.

Critical findings:

None

Major findings:

None

Minor findings:

None

COMMENTS:

Results are valid and can be used for decision making purposes.

Reviewers Name:

Rafael Infante

Chemist License 188

Signature:

Date:

April 16, 2016

SAMPLE ORGANIC DATA SAMPLE SUMMARY

Sample ID: JC16038-1

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.0	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.0	ug/L	1	-	υ	Yes
2,4-Dichlorophenol	2.0	ug/L	1	-	U	Yes
2,4-Dimethylphenol	5.0	ug/L	1	-	U	Yes
2,4-Dinitrophenol	10	ug/L	1	-	U	Yes
4,6-Dinitro-o-cresol	5.0	ug/L	1	-	U	Yes
2-Methylphenol	2.0	ug/L	1	-	U	Yes
3&4-Methylphenol	2.0	ug/L	1	-	U	Yes
2-Nitrophenol	5.0	ug/L	1	-	U	Yes
4-Nitrophenol	10	ug/L	1	~	U	Yes
Pentachlorophenol	5.0	ug/L	1	-	U	Yes
Phenol	2.0	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.0	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.0	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.0	ug/L	1	-	U	Yes
Acenaphthene	1.0	ug/L	1	-	U	Yes
Acenaphthylene	1.0	ug/L	1	-	U	Yes
Acetophenone	2.0	ug/L	1	~	U	Yes
Anthracene	1.0	ug/L	1	-	U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	5.0	ug/L	1	-	U	Yes
Benzo(a)anthracene	1.0	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.0	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.0	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.0	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.0	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.0	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.0	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.0	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.0	ug/L	1	-	U	Yes
4-Chloroaniline	1.3	ug/L	1	J	UJ	Yes
Carbazole	1.0	ug/L	1	-	U	Yes
Caprolactam	2.0	ug/L	1	-	U	Yes
Chrysene	1.0	ug/L	1	-	U	Yes
bis (2-Chloroethoxy) methane	2.0	ug/L	1	-	U	Yes
bis (2-Chloroethyl) ether	2.0	ug/L	1	(7)	U	Yes

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
bis(2-Chloroisopropyl)ether	2.0	ug/L	1	-	U	Yes
4-Chlorophenyl phenyl ether	2.0	ug/L	1	-	U	Yes
2,4-Dinitrotoluene	1.0	ug/L	1	-	U	Yes
2,6-Dinitrotoluene	1.0	ug/L	1	-	U	Yes
3,3'-Dichlorobenzidine	2.0	ug/L	1	-	U	Yes
Dibenzo(a,h)anthracene	1.0	ug/L	1	-	U	Yes
Dibenzofuran	5.0	ug/L	1	-	U	Yes
Di-n-butyl phthalate	2.0	ug/L	1	-	U	Yes
Di-n-octyl phthalate	2.0	ug/L	1	•	U	Yes
Diethyl phthalate	2.0	ug/L	1	-	U	Yes
Dimethyl phthalate	2.0	ug/L	1	-	U	Yes
bis(2-Ethylhexyl)phthalate	2.0	ug/L	1	-	υ	Yes
Fluoranthene	1.0	ug/L	1	-	U	Yes
Fluorene	1.0	ug/L	1	-	U	Yes
Hexachlorobenzene	1.0	ug/L	1	-	U	Yes
Hexachlorobutadiene	1.0	ug/L	1	-	U	Yes
Hexachlorocyclopentadiene	10	ug/L	1	-	U	Yes
Hexachloroethane	2.0	ug/L	1	-	U	Yes
Indeno(1,2,3-cd)pyrene	1.0	ug/L	1	-	U	Yes
Isophorone	2.0	ug/L	1	-	U	Yes
1-Methylnaphthalene	1.0	ug/L	1	-	U	Yes
2-Methylnaphthalene	1.0	ug/L	1	-	U	Yes
2-Nitroaniline	5.0	ug/L	1	-	U	Yes
3-Nitroaniline	5.0	ug/L	1	-	U	Yes
4-Nitroaniline	5.0	ug/L	1	-	U	Yes
Naphthalene	1.0	ug/L	1	-	U	Yes
Nitrobenzene	2.0	ug/L	1	-	U	Yes
N-Nitroso-di-n-propylamine	2.0	ug/L	1	-	U	Yes
Nitrosodiphenylamine	5.0	ug/L	1	-	U	Yes
Phenanthrene	1.0	ug/L	1	-	U	Yes
Pyrene	1.0	ug/L	1	-	U	Yes
1,2,4,5-Tetrachlorobenzene	2.0	ug/L	1	-	U	Yes
METHOD:	8270D (S	IM)				
Naphthalene	0.10	ug/L	1	_	U	Yes
1,4-Dioxane	0.10	ug/L	1	-	Ü	Yes

.

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-2

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.3	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.3	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.1	ug/L	1	-	U	Yes
2,4-Dimethylphenol	5.3	ug/L	1	-	U	Yes
2,4-Dinitrophenol	11	ug/L	1	-	U	Yes
4,6-Dinitro-o-cresol	5.3	ug/L	1	-	U	Yes
2-Methylphenol	2.1	ug/L	1	-	U	Yes
3&4-Methylphenoi	2.1	ug/L	1	-	U	Yes
2-Nitrophenol	5.3	ug/L	1	-	U	Yes
4-Nitrophenol	11	ug/L	1	-	U	Yes
Pentachlorophenol	5.3	ug/L	1	-	U	Yes
Phenol	2.1	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.3	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.3	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.3	ug/L	1	-	U	Yes
Acenaphthene	1.1	ug/L	1	-	U	Yes
Acenaphthylene	1.1	ug/L	1	-	U	Yes
Acetophenone	2.1	ug/L	1	-	U	Yes
Anthracene	1.1	ug/L	1		U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	5.3	ug/L	1	-	U	Yes
Benzo(a)anthracene	1.1	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.1	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.1	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.1	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.1	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.1	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.1	ug/L	1	-	Ü	Yes
1,1'-Biphenyl	1.1	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.1	ug/L	1	-	U	Yes
4-Chloroaniline	5.3	ug/L	1	-	U	Yes
Carbazole	1.1	ug/L	1	-	U	Yes
Caprolactam	2.1	ug/L	1	-	U	Yes
Chrysene	1.1	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.1	ug/L	1	-	U	Yes
bis(2-Chloroethyl)ether	2.1	ug/L	1	-	U	Yes

.

METHOD:	02/UD					
Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
bis (2-Chlorois opropy!) ether	2.1	ug/L	1	-	U	Yes
4-Chlorophenyl phenyl ether	2.1	ug/L	1	-	Ų	Yes
2,4-Dinitrotoluene	1.1	ug/L	1	-	U	Yes
2,6-Dînitrotoluene	1.1	ug/L	1	-	U	Yes
3,3'-Dichlorobenzidine	2.1	ug/L	1	-	U	Yes
Dibenzo(a,h)anthracene	1.0	ug/L	1	-	U	Yes
Dibenzofuran	5.3	ug/L	1	-	U	Yes
Di-n-butyl phthalate	2.1	ug/L	1	-	U	Yes
Di-n-octyl phthalate	2.1	ug/L	1	-	U	Yes
Diethyl phthalate	2.1	ug/L	1	-	U	Yes
Dimethyl phthalate	2.1	ug/L	1	-	U	Yes
bis(2-Ethylhexyl)phthalate	2.1	ug/L	1	-	U	Yes
Fluoranthene	1.1	ug/L	1	-	U	Yes
Fluorene	1.1	ug/L	1	-	U	Yes
Hexachlorobenzene	1.1	ug/L	1	-	U	Yes
Hexachlorobutadiene	1.1	ug/L	1	-	U	Yes
Hexachlorocyclopentadiene	11	ug/L	1	~	U	Yes
Hexachloroethane	2.1	ug/L	1	-	U	Yes
Indeno(1,2,3-cd)pyrene	1.1	ug/L	1	-	U	Yes
Isophorone	2.1	ug/L	1	-	U	Yes
1-Methylnaphthalene	1.1	ug/L	1	-	U	Yes
2-Methylnaphthalene	1.1	ug/L	1	-	U	Yes
2-Nitroaniline	5.3	ug/L	1	-	U	Yes
3-Nitroaniline	5.3	ug/L	1	-	U	Yes
4-Nitroaniline	5.3	ug/L	1	-	U	Yes
Naphthalene	1.1	ug/L	1	-	U	Yes
Nitrobenzene	2.1	ug/L	1	-	U	Yes
N-Nitroso-di-n-propylamine	2.1	ug/L	1	-	U	Yes
Nitrosodiphenylamine	5.3	ug/L	1	-	U	Yes
Phenanthrene	1.1	ug/L	1	-	U	Yes
Pyrene	1.1	ug/L	1	-	U	Yes
1,2,4,5-Tetrachlorobenzene	2.1	ug/L	1	-	U	Yes
METHOD: 8	32 70D (S	IM)				
Naphthalene	0.11	ug/L	1	-	U	Yes
1,4-Dioxane	1.72	ug/L	1	-	-	Yes

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-3

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016

Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.0	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.0	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.0	ug/L	1	-	U	Yes
2,4-Dimethylphenol	5.0	ug/L	1	-	U	Yes
2,4-Dinitrophenol	10	ug/L	1	-	U	Yes
4,6-Dinitro-o-cresol	5.0	ug/L	1	-	U	Yes
2-Methylphenol	2.0	ug/L	1	-	U	Yes
3&4-Methylphenol	2.0	ug/L	1	-	U	Yes
2-Nitrophenol	5.0	ug/L	1	-	U	Yes
4-Nitrophenol	10	ug/L	1	-	U	Yes
Pentachlorophenol	5.0	ug/L	1	-	U	Yes
Phenol	2.0	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.0	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.0	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.0	ug/L	1	-	U	Yes
Acenaphthene	0.67	ug/L	1	J	UJ	Yes
Acenaphthylene	1.0	ug/L	1	-	U	Yes
Acetophenone	2.0	ug/L	1	-	U	Yes
Anthracene	1.0	ug/L	1	-	U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	5.0	ug/L	1	-	U	Yes
Benzo(a) anthracene	1.0	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.0	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.0	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.0	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.0	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.0	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.0	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.0	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.0	ug/L	1	-	U	Yes
4-Chloroaniline	5.0	ug/L	1	-	U	Yes
Carbazole	1.0	ug/L	1	-	U	Yes
Caprolactam	2.0	ug/L	1	-	U	Yes
Chrysene	1.0	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.0	ug/L	1	-	U	Yes
bis(2-Chloroethyl)ether	2.0	ug/L	1	-	U	Yes

Analyta Nama		1114	Dilukias Frak	Labori.	Mandada et a	0
Analyte Name	Result		Dilution Factor	Lab Flag		-
bis(2-Chloroisopropyl)ether	2.0	ug/L	1	*	U	Yes
4-Chlorophenyl phenyl ether	2.0	ug/L	1	-	U	Yes
2,4-Dinitrotoluene	1.0	ug/L	1	-	U	Yes
2,6-Dinitrotoluene	1.0	ug/L	1	-	U ·	Yes
3,3'-Dichlorobenzidine	2.0	ug/L	1	-	U	Yes
Dibenzo(a,h)anthracene	1.0	ug/L	1	-	U	Yes
Dibenzofuran	5.0	ug/L	1	-	U	Yes
Di-n-butyl phthalate	2.0	ug/L	1	-	U	Yes
Di-n-octyl phthalate	2.0	ug/L	1	-	U	Yes
Diethyl phthalate	2.0	ug/L	1	-	U	Yes
Dimethyl phthalate	2.0	ug/L	1	-	U	Yes
bis(2-Ethylhexyl)phthalate	2.0	ug/L	1	-	U	Yes
Fluoranthene	1.0	ug/L	1	-	U	Yes
Fluorene	1.2	ug/L	1	-	-	Yes
Hexachlorobenzene	1.0	ug/L	1	-	U	Yes
Hexachlorobutadiene	1.0	ug/L	1	-	U	Yes
Hexachlorocyclopentadiene	10	ug/L	1	-	U	Yes
Hexachloroethane	2.0	ug/L	1	-	U	Yes
Indeno(1,2,3-cd)pyrene	1.0	ug/L	1	-	U	Yes
Isophorone	2.0	ug/L	1	-	U	Yes
1-Methylnaphthalene	1.0	ug/L	1	-	U	Yes
2-Methylnaphthalene	52.4	ug/L	1	-	-	Yes
2-Nitroaniline	5.0	ug/L	1	-	U	Yes
3-Nitroaniline	5.0	ug/L	1	-	U	Yes
4-Nitroaniline	5.0	ug/L	1	-	U	Yes
Naphthalene	3.6	ug/L	1	-	-	Yes
Nitrobenzene	2.0	ug/L	1	-	U	Yes
N-Nitroso-di-n-propylamine	2.0	ug/L	1	-	U	Yes
Nitrosodiphenylamine	5.0	ug/L	1	-	U	Yes
Phenanthrene	1.0	ug/L	1	-	U	Yes
Pyrene	1.0	ug/L	1	_	U	Yes
1,2,4,5-Tetrachiorobenzene	2.0	ug/L	1	-	U	Yes
		<u> </u>				
METHOD:	8270D (S	IM)				
Naphthalene	3.57	ug/L	1	-	-	Yes
1,4-Dioxane	0.10	ug/L	1	-	U	Yes

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-4

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.3	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.3	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.1	ug/L	1	-	U	Yes
2,4-Dimethylphenol	5.7	ug/L	1	-	-	Yes
2,4-Dinitrophenol	11	ug/L	1	-	U	Yes
4,6-Dinitro-o-cresol	5.3	ug/L	1	-	U	Yes
2-Methylphenol	2.1	ug/L	1	-	U	Yes
3&4-Methylphenol	2.1	ug/L	1	-	U	Yes
2-Nitrophenol	5.3	ug/L	1	-	U	Yes
4-Nitrophenol	11	ug/L	1	-	U	Yes
Pentachlorophenol	5.3	ug/L	1	-	U	Yes
Phenol	2.1	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.3	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.3	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.3	ug/L	1	-	U	Yes
Acenaphthene	1.1	ug/L	1	-	U	Yes
Acenaphthylene	1.1	ug/L	1	-	U	Yes
Acetophenone	2.1	ug/L	1	-	U	Yes
Anthracene	1.1	ug/L	1	-	U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	5.3	ug/L	1	-	U	Yes
Benzo(a)anthracene	1.1	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.1	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.1	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.1	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.1	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.1	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.1	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.1	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.1	ug/L	1	-	U	Yes
4-Chloroaniline	5.3	ug/L	1	-	U	Yes
Carbazole	1.1	ug/L	1	-	U	Yes
Caprolactam	2.1	ug/L	1	-	U	Yes
Chrysene	1.1	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.1	ug/L	1	-	U	Yes
bis(2-Chloroethyl)ether	2.1	ug/L	1	-	U	Yes

WETHOD:						
Analyte Name	Result		Dilution Factor	Lab Flag		Reportable
bis(2-Chloroisopropyl)ether	2.1	ug/L	1	-	U	Yes
4-Chlorophenyl phenyl ether	2.1	ug/L	1	-	U	Yes
2,4-Dinitrotoluene	1.1	ug/L	1	-	U	Yes
2,6-Dinitrotoluene	1.1	ug/L	1	-	U	Yes
3,3'-Dichlorobenzidine	2.1	ug/L	1	-	U	Yes
1,4-Dioxane	1.0	ug/L	1	-	U	Yes
Dibenzo(a,h)anthracene	1.1	ug/L	1	-	U	Yes
Dibenzofuran	5.3	ug/L	1	-	U	Yes
Di-n-butyl phthalate	2.1	ug/L	1	-	Ų	Yes
Di-n-octyl phthalate	2.1	ug/L	1	-	U	Yes
Diethyl phthalate	2.1	ug/L	1	-	U	Yes
Dimethyl phthalate	2.1	ug/L	1	-	U	Yes
bis(2-Ethylhexyl)phthalate	2.1	ug/L	1	-	U	Yes
Fluoranthene	1.1	ug/L	1	-	U	Yes
Fluorene	1.1	ug/L	1	-	U	Yes
Hexachlorobenzene	1.1	ug/L	1	-	U	Yes
Hexachlorobutadiene	1.1	ug/L	1	-	U	Yes
Hexachlorocyclopentadiene	11	ug/L	1	-	U	Yes
Hexachloroethane	2.1	ug/L	1	-	U	Yes
Indeno(1,2,3-cd)pyrene	1.1	ug/L	1	-	U	Yes
Isophorone	2.1	ug/L	1	-	U	Yes
1-Methylnaphthalene	1.1	ug/L	1	-	U	Yes
2-Methylnaphthalene	0.63	ug/L	1	J	UJ	Yes
2-Nitroaniline	5.3	ug/L	1	-	U	Yes
3-Nitroaniline	5.3	ug/L	1	-	U	Yes
4-Nitroaniline	1.1	ug/L	1	-	U	Yes
Naphthalene	2.7	ug/L	1	-	-	Yes
Nitrobenzene	2.1	ug/L	1	-	U	Yes
N-Nitroso-di-n-propylamine	2.1	ug/L	1	-	U	Yes
Nitrosodiphenylamine	5.3	ug/L	1	-	U	Yes
Phenanthrene	1.1	ug/L	1	-	U	Yes
Pyrene	1.1	ug/L	1	-	U	Yes
1,2,4,5-Tetrachlorobenzene	2.1	ug/L	1	-	U	Yes
METHOD:	9270D /C	(6.41)				
Naphthalene	2.48	ug/L	1	_		Yes
1,4-Dioxane	0.734	-		-	-	
T'4-DIOXQUE	U./54	ug/L	1	-	-	Yes

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-5

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.3	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.3	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.1	ug/L	1	-	Ų	Yes
2,4-Dimethylphenol	5.3	ug/L	1	-	U	Yes
2,4-Dinitrophenol	11	ug/L	1	-	Ų	Yes
4,6-Dinitro-o-cresol	5.3	ug/L	1	-	U	Yes
2-Methylphenol	2.1	ug/L	1	-	U	Yes
3&4-Methylphenol	2.1	ug/L	1	-	U	Yes
2-Nitrophenol	5.3	ug/L	1	-	U	Yes
4-Nitrophenol	11	ug/L	1	-	U	Yes
Pentachlorophenol	5.3	ug/L	1	-	U	Yes
Phenol	2.1	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.3	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.3	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.3	ug/L	1	-	U	Yes
Acenaphthene	1.1	ug/L	1	-	U	Yes
Acenaphthylene	1.1	ug/L	1	-	U	Yes
Acetophenone	2.1	ug/L	1	-	U	Yes
Anthracene	1.1	ug/L	1	-	U	Yes
Atrazine	1.1	ug/L	1	-	U	Yes
Benzaldehyde	5.3	ug/L	1	-	U	Yes
Benzo(a)anthracene	1.1	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.1	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.1	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.1	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.1	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.1	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.1	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.1	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.1	ug/L	1	-	U	Yes
4-Chloroaniline	5.3	ug/L	1	-	U	Yes
Carbazole	1.1	ug/L	1	-	U	Yes
Caprolactam	2.1	ug/L	1	-	U	Yes
Chrysene	1.1	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.1	ug/L	1	-	U	Yes
bis(2-Chloroethyl)ether	2.1	ug/L	1	-	U	Yes

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable			
bis(2-Chloroisopropyl)ether	2.1	ug/L	1	-	U	Yes			
4-Chlorophenyl phenyl ether	2.1	ug/L	1	_	Ū	Yes			
2,4-Dinitrotoluene	1.1	ug/L	1	_	Ū	Yes			
2,6-Dinitrotoluene	1.1	ug/L	1	_	Ü	Yes			
3,3'-Dichlorobenzidine	2.1	ug/L	1	_	Ū	Yes			
Dibenzo(a,h)anthracene	1.1	ug/L	1	-	U	Yes			
Dibenzofuran	1.1	ug/L	1	-	U	Yes			
Di-n-butyl phthalate	5.3	ug/L	1	-	U	Yes			
Di-n-octyl phthalate	2.1	ug/L	1	-	U	Yes			
Diethyl phthalate	2.1	ug/L	1	-	U	Yes			
Dimethyl phthalate	2.1	ug/L	1	_	U	Yes			
bis(2-Ethylhexyl)phthalate	2.1	ug/L	1	-	U	Yes			
Fluoranthene	2.1	ug/L	1	-	U	Yes			
Fluorene	1.1	ug/L	1	-	U	Yes			
Hexachlorobenzene	1.1	ug/L	1	-	U	Yes			
Hexachlorobutadiene	1.1	ug/L	1	-	U	Yes			
Hexachlorocyclopentadiene	1.1	ug/L	1	_	U	Yes			
Hexachloroethane	11	ug/L	1	-	U	Yes			
Indeno(1,2,3-cd)pyrene	2.1	ug/L	1	-	U	Yes			
Isophorone	1.1	ug/L	1	-	U	Yes			
1-Methylnaphthalene	2.1	ug/L	1	-	U	Yes			
2-Methylnaphthalene	1.1	ug/L	1	-	J	Yes			
2-Nitroaniline	5.3	ug/L	1	-	U	Yes			
3-Nitroaniline	5.3	ug/L	1	-	U	Yes			
4-Nitroaniline	5.3	ug/L	1	-	U	Yes			
Naphthalene	1.1	ug/L	1	-	J	Yes			
Nitrobenzene	2.1	ug/L	1	-	U	Yes			
N-Nitroso-di-n-propylamine	2.1	ug/L	1	-	U	Yes			
Nitrosodiphenylamine	5.3	ug/L	1	-	U	Yes			
Phenanthrene	1.1	ug/L	1	-	U	Yes			
Pyrene	1.1	ug/L	1	-	U	Yes			
1,2,4,5-Tetrachlorobenzene	2.1	ug/L	1	-	U	Yes			
METHOD: 8270D (SIM)									
	-	•	4			Ves			
Naphthalene	0.11	ug/L	1	-	-	Yes			
1,4-Dioxane	0.388	ug/L	1	-	-	Yes			

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-6

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.2	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.2	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.1	ug/L	1	-	U	Yes
2,4-Dimethylphenol	5.3	ug/L	1	-	U	Yes
2,4-Dinitrophenol	11	ug/L	1	-	U	Yes
4,6-Dinitro-o-cresol	5.2	ug/L	1	-	U	Yes
2-Methylphenol	2.1	ug/L	1	-	U	Yes
3&4-Methylphenol	2.1	ug/L	1	7.	U	Yes
2-Nitrophenol	5.2	ug/L	1	-	U	Yes
4-Nitrophenol	10	ug/L	1	-	U	Yes
Pentachlorophenol	5.2	ug/L	1	07	U	Yes
Phenol	2.1	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.2	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.2	ug/L	1	0.45	U	Yes
2,4,6-Trichlorophenol	5.2	ug/L	1	-	U	Yes
Acenaphthene	0.44	ug/L	1	J	UJ	Yes
Acenaphthylene	1.0	ug/L	1	-	U	Yes
Acetophenone	2.1	ug/L	1	5.6	U	Yes
Anthracene	1.0	ug/L	1	-	U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	5.2	ug/L	1	-	U	Yes
Benzo(a)anthracene	1.0	ug/L	1	0.20	U	Yes
Benzo(a)pyrene	1.0	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.0	ug/L	1		U	Yes
Benzo(g,h,i)perylene	1.0	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.0	ug/L	1		U	Yes
4-Bromophenyl phenyl ether	2.1	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.1	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.1	ug/L	1	6.70	U	Yes
2-Chloronaphthalene	2.1	ug/L	1	_	U	Yes
4-Chloroaniline	5.2	ug/L	1	-	U	Yes
Carbazole	1.0	ug/L	1	47	U	Yes
Caprolactam	2.1	ug/L	1	-	U	Yes
Chrysene	1.0	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.1	ug/L	1	-	U	Yes
bis (2-Chloroethyl) ether	2.1	ug/L	1	+	U	Yes

WEITOD.										
Analyte Name	Result		Dilution Factor	Lab Flag		•				
bis(2-Chloroisopropyl)ether	2.1	ug/L	1	-	U	Yes				
4-Chlorophenyl phenyl ether	2.1	ug/L	1	-	U	Yes				
2,4-Dinitrotoluene	1.0	ug/L	1	-	U	Yes				
2,6-Dinitrotoluene	1.0	ug/L	1	-	U	Yes				
3,3'-Dichlorobenzidine	2.1	ug/L	1	-	U	Yes				
Dibenzo(a,h)anthracene	1.0	ug/L	1	-	U	Yes				
Dibenzofuran	1.0	ug/L	1	-	U	Yes				
Di-n-butyl phthalate	5.2	ug/L	1	-	U	Yes				
Di-n-octyl phthalate	2.1	ug/L	1	-	U	Yes				
Diethyl phthalate	2.1	ug/L	1	-	U	Yes				
Dimethyl phthalate	2.1	ug/L	1	-	U	Yes				
bis (2-Ethylhexyl) phthalate	2.1	ug/L	1	-	U	Yes				
Fluoranthene	2.1	ug/L	1	-	U	Yes				
Fluorene	0.49	ug/L	1	J	UJ	Yes				
Hexachlorobenzene	1.0	ug/L	1	-	U	Yes				
Hexachlorobutadiene	1.0	ug/L	1	-	U	Yes				
Hexachlorocyclopentadiene	1.1	ug/L	1	-	U	Yes				
Hexachloroethane	10	ug/L	1	-	U	Yes				
Indeno(1,2,3-cd)pyrene	2.1	ug/L	1	-	U	Yes				
Isophorone	1.0	ug/L	1	-	U	Yes				
1-Methylnaphthalene	2.1	ug/L	1	-	U	Yes				
2-Methylnaphthalene	3.4	ug/L	1	-	J	Yes				
2-Nitroaniline	5.2	ug/L	1	-	U	Yes				
3-Nitroaniline	5.2	ug/L	1	-	U	Yes				
4-Nitroaniline	5.2	ug/L	1	-	U	Yes				
Naphthalene	2.3	ug/L	1	-	J	Yes				
Nitrobenzene	2.1	ug/L	1	-	Ų	Yes				
N-Nitroso-di-n-propylamine	2.1	ug/L	1	-	U	Yes				
Nitrosodiphenylamine	5.2	ug/L	1	-	U	Yes				
Phenanthrene	0.61	ug/L	1	J	UJ	Yes				
Pyrene	1.0	ug/L	1	-	U	Yes				
1,2,4,5-Tetrachlorobenzene	2.1	ug/L	1	-	U	Yes				
METHOD: 8270D (SIM)										
Naphthalene	0.11	ug/L	1	-	-	Yes				
1,4-Dioxane	0.388	ug/L	1	-	-	Yes				

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-8

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.1	ug/L	1		U	Yes
4-Chloro-3-methyl phenol	5.1	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.0	ug/L	1	-	U	Yes
2,4-Dimethylphenol	5.1	ug/L	1	-	U	Yes
2,4-Dinitrophenol	10	ug/L	1	_	U	Yes
4,6-Dinitro-o-cresol	5.1	ug/L	1	190	U	Yes
2-Methylphenol	2.0	ug/L	1	15.0	U	Yes
3&4-Methylphenol	2.0	ug/L	1	_	U	Yes
2-Nitrophenol	5.1	ug/L	1	(+)	U	Yes
4-Nitrophenol	10	ug/L	1	-	U	Yes
Pentachlorophenol	5.1	ug/L	1	-	U	Yes
Phenol	2.0	ug/L	1		U	Yes
2,3,4,6-Tetrachlorophenol	5.1	ug/L	1	_	U	Yes
2,4,5-Trichlorophenol	5.1	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.1	ug/L	1	-	U	Yes
Acenaphthene	1.0	ug/L	1	-	U	Yes
Acenaphthylene	1.0	ug/L	1	-	U	Yes
Acetophenone	2.0	ug/L	1	-	U	Yes
Anthracene	1.0	ug/L	1	-	U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	0.90	ug/L	1	j	UJ	Yes
Benzo(a)anthracene	1.0	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.0	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.0	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.0	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.0	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.0	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.0	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.0	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.0	ug/L	1	-	U	Yes
4-Chloroaniline	5.1	ug/L	1	-	U	Yes
Carbazole	1.0	ug/L	1	-	U	Yes
Caprolactam	2.0	ug/L	1	-	U	Yes
Chrysene	1.0	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.0	ug/L	1	-	U	Yes
bis(2-Chloroethyl)ether	2.0	ug/L	1	-	U	Yes

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
bis(2-Chloroisopropyl)ether	2.0	ug/L	1		U	Yes
4-Chlorophenyl phenyl ether	2.0	ug/L	1	-	Ü	Yes
2,4-Dinitrotoluene	1.0	ug/L	1	-	U	Yes
2,6-Dinitrotoluene	1.0	ug/L	1	_	U	Yes
3,3'-Dichlorobenzidine	2.0	ug/L	1	-	U	Yes
1,4-Dioxane	2220	ug/L	40		-	Yes
Dibenzo(a,h)anthracene	1.0	ug/L	1	_	U	Yes
Dibenzofuran	5.1	ug/L	1		U	Yes
Di-n-butyl phthalate	2.0	ug/L	1	-	U	Yes
Di-n-octyl phthalate	2.0	ug/L	1	140	U	Yes
Diethyl phthalate	2.0	ug/L	1	-	U	Yes
Dimethyl phthalate	2.0	ug/L	1	12	U	Yes
bis(2-Ethylhexyl)phthalate	2.0	ug/L	1	-	U	Yes
Fluoranthene	1.0	ug/L	1		U	Yes
Fluorene	1.0	ug/L	1		U	Yes
Hexachlorobenzene	1.0	ug/L	1	-	U	Yes
Hexachlorobutadiene	1.0	ug/L	1	-	U	Yes
Hexachlorocyclopentadiene	10	ug/L	1	-	U	Yes
Hexachloroethane	2.0	ug/L	1	-	U	Yes
Indeno(1,2,3-cd)pyrene	1.0	ug/L	1	02.0	U	Yes
Isophorone	2.0	ug/L	1	-	U	Yes
1-Methylnaphthalene	1.0	ug/L	1	-	U	Yes
2-Methylnaphthalene	1.0	ug/L	1	-	U	Yes
2-Nitroaniline	5.1	ug/L	1	-	U	Yes
3-Nitroaniline	5.1	ug/L	1	-	U	Yes
4-Nitroaniline	5.1	ug/L	1	-	U	Yes
Naphthalene	1.0	ug/L	1	-	U	Yes
Nitrobenzene	2.0	ug/L	1	-	U	Yes
N-Nitroso-di-n-propylamine	2.0	ug/L	1	-	U	Yes
Nitrosodiphenylamine	5.1	ug/L	1	-	U	Yes
Phenanthrene	1.0	ug/L	1	-	U	Yes
Pyrene	1.0	ug/L	1	-	U	Yes
1,2,4,5-Tetrachlorobenzene	2.0	ug/L	1	-	U	Yes
METHOD:	8270D (S	IM)				
Naphthalene	0.10	ug/L	1	-	U	Yes

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-9

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.0	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.0	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.0	ug/L	1	-	U	Yes
2,4-Dimethylphenol	5.0	ug/L	1	-	U	Yes
2,4-Dinitrophenol	10	ug/L	1	-	Ų	Yes
4,6-Dinitro-o-cresol	5.0	ug/L	1	-	Ų	Yes
2-Methylphenol	2.0	ug/L	1	-	U	Yes
3&4-Methylphenol	2.0	ug/L	1	-	U	Yes
2-Nitrophenol	5.0	ug/L	1	-	U	Yes
4-Nitrophenol	10	ug/L	1	-	U	Yes
Pentachlorophenol	5.0	ug/L	1	-	U	Yes
Phenol	2.0	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.0	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.0	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.0	ug/L	1	-	U	Yes
Acenaphthene	1.0	ug/L	1	-	U	Yes
Acenaphthylene	1.0	ug/L	1	-	U	Yes
Acetophenone	2.0	ug/L	1	-	U	Yes
Anthracene	1.0	ug/L	1	-	U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	5.0	ug/L	1	-	U	Yes
Benzo(a)anthracene	1.0	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.0	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.0	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.0	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.0	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.0	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.0	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.0	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.0	ug/L	1	-	U	Yes
4-Chloroaniline	5.0	ug/L	1	-	UJ	Yes
Carbazole	1.0	ug/L	1	-	U	Yes
Caprolactam	2.0	ug/L	1	-	U	Yes
Chrysene	1.0	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.0	ug/L	1	-	U	Yes
bis (2-Chloroethyl) ether	2.0	ug/L	1	-	U	Yes

	WIETTIOD: 1	52700								
	Analyte Name	Result		Dilution Factor	Lab Flag		•			
	bis(2-Chloroisopropyl)ether	2.0	ug/L	1	-	U	Yes			
	4-Chlorophenyl phenyl ether	2.0	ug/L	1 0	-	U	Yes			
	2,4-Dinitrotoluene	1.0	ug/L	1	-	U	Yes			
	2,6-Dinitrotoluene	1.0	ug/L	1	-	U	Yes			
	3,3'-Dichlorobenzidine	2.0	ug/L	1	-	U	Yes			
	Dibenzo(a,h)anthracene	1.0	ug/L	1	-	U	Yes			
	Dibenzofuran	5.0	ug/L	1	-	U	Yes			
	Di-n-butyl phthalate	2.0	ug/L	1	-	U	Yes			
	Di-n-octyl phthalate	2.0	ug/L	1	-	U	Yes			
	Diethyl phthalate	2.0	ug/L	1	-	U	Yes			
	Dimethyl phthalate	2.0	ug/L	1	-	U	Yes			
	bis(2-Ethylhexyl)phthalate	1.2	ug/L	1	J	UJ	Yes			
	Fluoranthene	1.0	ug/L	1	-	U	Yes			
	Fluorene	1.0	ug/L	1	-	U	Yes			
	Hexachlorobenzene	1.0	ug/L	1	_	U	Yes			
	Hexachlorobutadiene	1.0	ug/L	1	-	U	Yes			
	Hexachlorocyclopentadiene	10	ug/L	1	-	U	Yes			
	Hexachloroethane	2.0	ug/L	1	•	U	Yes			
	Indeno(1,2,3-cd)pyrene	1.0	ug/L	1	-	U	Yes			
	Isophorone	2.0	ug/L	1	-	U	Yes			
	1-Methylnaphthalene	1.0	ug/L	1	•	U	Yes			
	2-Methylnaphthalene	1.0	ug/L	1	-	U	Yes			
	2-Nitroaniline	5.0	ug/L	1	-	U	Yes			
	3-Nitroaniline	5.0	ug/L	1	-	U	Yes			
	4-Nitroaniline	5.0	ug/L	1	-	U	Yes			
	Naphthalene	1.0	ug/L	1	-	U	Yes			
	Nitrobenzene	2.0	ug/L	1	-	U	Yes			
	N-Nitroso-di-n-propylamine	2.0	ug/L	1	-	U	Yes			
	Nitrosodiphenylamine	5.0	ug/L	1	-	U	Yes			
	Phenanthrene	1.0	ug/L	1	_	U	Yes			
	Pyrene	1.0	ug/L	1	-	U	Yes			
	1,2,4,5-Tetrachlorobenzene	2.0	ug/L	1	-	U	Yes			
	METHOD: 8270D (SIM)									
	Naphthalene	0.10	ug/L	1	-	U	Yes			
	1,4-Dioxane	0.10	ug/L	1	-	U	Yes			

Analyte Name Result Units Dilution Factor Lab Flag Validation Reportable

Sample ID: JC16038-10

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
2-Chlorophenol	5.2	ug/L	1	-	U	Yes
4-Chloro-3-methyl phenol	5.2	ug/L	1	-	U	Yes
2,4-Dichlorophenol	2.1	ug/L	1	12	U	Yes
2,4-Dimethylphenol	5.3	ug/L	1	-	Ų	Yes
2,4-Dinitrophenol	11	ug/L	1	.73	U	Yes
4,6-Dinitro-o-cresol	5.2	ug/L	1	-	U	Yes
2-Methylphenol	2.1	ug/L	1	-	U	Yes
3&4-Methylphenol	2.1	ug/L	1	-	U	Yes
2-Nitrophenol	5.2	ug/L	1	-	U	Yes
4-Nitrophenol	10	ug/L	1	-	U	Yes
Pentachlorophenol	5.2	ug/L	1	-	U	Yes
Phenol	2.1	ug/L	1	-	U	Yes
2,3,4,6-Tetrachlorophenol	5.2	ug/L	1	-	U	Yes
2,4,5-Trichlorophenol	5.2	ug/L	1	-	U	Yes
2,4,6-Trichlorophenol	5.2	ug/L	1	-	U	Yes
Acenaphthene	1.0	ug/L	1	-	U	Yes
Acenaphthylene	1.0	ug/L	1	~	U	Yes
Acetophenone	2.1	ug/L	1	-	U	Yes
Anthracene	1.0	ug/L	1	-	U	Yes
Atrazine	1.0	ug/L	1	-	U	Yes
Benzaldehyde	5.2	ug/L	1	-	U	Yes
Benzo(a)anthracene	1.0	ug/L	1	-	U	Yes
Benzo(a)pyrene	1.0	ug/L	1	-	U	Yes
Benzo(b)fluoranthene	1.0	ug/L	1	-	U	Yes
Benzo(g,h,i)perylene	1.0	ug/L	1	-	U	Yes
Benzo(k)fluoranthene	1.0	ug/L	1	-	U	Yes
4-Bromophenyl phenyl ether	2.1	ug/L	1	-	U	Yes
Butyl benzyl phthalate	2.1	ug/L	1	-	U	Yes
1,1'-Biphenyl	1.1	ug/L	1	-	U	Yes
2-Chloronaphthalene	2.1	ug/L	1	-	U	Yes
4-Chloroaniline	5.2	ug/L	1	-	U	Yes
Carbazole	1.0	ug/L	1	-	U	Yes
Caprolactam	2.1	ug/L	1	-	U	Yes
Chrysene	1.0	ug/L	1	-	U	Yes
bis(2-Chloroethoxy)methane	2.1	ug/L	1	-	U	Yes
bis (2-Chloroethyl) ether	2.1	ug/L	1	-	U	Yes

Analyte Name	Result	Unito	Dilution Factor	Lah Elaa	Validation	Panartabla
bis(2-Chloroisopropyl)ether	2.1	ug/L	1	ran Liag	U	Yes
4-Chlorophenyl phenyl ether	2.1	ug/L	1	-	U	Yes
2,4-Dinitrotoluene	1.0	ug/L	1	-	U	Yes
2,6-Dinitrotoluene	1.0	ug/L	1	-	U	Yes
3,3'-Dichlorobenzidine	2.1	ug/L	1	_	U	Yes
1,4-Dioxane	385	ug/L	5	-	-	Yes
Dibenzo(a,h)anthracene	1.0	ug/L	1	-	- U	Yes
Dibenzofuran	1.0	ug/L	1	_	U	Yes
Di-n-butyl phthalate	5.2	ug/L	1	_	U	Yes
Di-n-octyl phthalate	2.1	ug/L	1	-	U	Yes
Diethyl phthalate	2.1	ug/L	1	_	U	Yes
Dimethyl phthalate	2.1	ug/L	1	_	U	Yes
bis(2-Ethylhexyl)phthalate	2.1	ug/L	1	_	U	Yes
Fluoranthene	2.1	ug/L	1	-	U	Yes
Fluorene	1.0	ug/L	1	J	ΩJ	Yes
Hexachlorobenzene	1.0	ug/L	1	_	Ü	Yes
Hexachlorobutadiene	1.0	ug/L	1	_	U	Yes
Hexachlorocyclopentadiene	10.0	ug/L	1	_	U	Yes
Hexachloroethane	2	ug/L	1	-	U	Yes
Indeno(1,2,3-cd)pyrene	1.0	ug/L	1	-	U	Yes
Isophorone	2.1	ug/L	1	•	U	Yes
2-Methylnaphthalene	1.0	ug/L	1	_	U	Yes
2-Nitroaniline	5.2	ug/L	1	-	U	Yes
3-Nitroaniline	5.2	ug/L	1	-	U	Yes
4-Nitroaniline	5.2	ug/L	1		U	Yes
Naphthalene	1.0	ug/L	1	_	U	Yes
Nitrobenzene	2.1	ug/L	1	_	U	Yes
N-Nitroso-di-n-propylamine	2.1	ug/L	1	-	U	Yes
Nitrosodiphenylamine	5.2	ug/L	1	_	U	Yes
Phenanthrene	1.0	ug/L	1	•	U	Yes
Pyrene	1.0	ug/L	1	_	Ü	Yes
1,2,4,5-Tetrachlorobenzene	2.1	ug/L	1	-	Ü	Yes
,, .,		-0/ -	476		-	
METHOD:	8270D (S	IM)				
Naphthalene	0.10	ug/L	1	-	-	Yes

	Project Number:_JC16038 Date:_March_8-9,_2016 Shipping Date:_March_9-10,_2016 EPA Region:2
REVIEW OF SEMIVOLATILE OR	GANIC PACKAGE
The following guidelines for evaluating volatile required validation actions. This document will assigned judgment to make more informed decision and in users. The sample results were assessed according documents in the following order of precedent Section, SOP HW-35A, July 2015 –Revision 0. Semivorand data validation actions listed on the data reviguidance document, unless otherwise noted.	sist the reviewer in using professional better serving the needs of the data of the data to USEPA data validation guidance se: EPA Hazardous Waste Supportulatile Data Validation. The QC criteria
The hardcopied (laboratory name) _Accutest	
Lab. Project/SDG No.:JC16038 No. of Samples:9_Full_scan/9_SIM	Sample matrix:Groundwater
Trip blank No.:	X Laboratory Control SpikesX Field DuplicatesX CalibrationsX Compound IdentificationsX Compound QuantitationX Quantitation Limits
Definition of Qualifiers:	
J- Estimated results U- Compound not detected R- Rejected data UJ- Estimated nondetect Reviewer: Rafuel Defaut	

DATA COMPLETENESS

MISSING INFORMATION	DATE LAB. CONTACTED	DATE RECEIVED
		
	'	
		<u> </u>
		<u> </u>

All criteria were met	X
Criteria were not mel	
and/or see below	_

HOLDING TIMES

The objective of this parameter is to ascertain the validity of the results based on the holding time of the sample from time of collection to the time of analysis.

Complete table for all samples and note the analysis and/or preservation not within criteria

SAMPLE ID	DATE SAMPLED	DATE EXTRACTED/ANALYZED	рН	ACTION
All samples extracted	d and analyzed wil	thin method recommended ho	lding 1	time.
	iii			

Cooler temperature (Criteria	: 4 <u>+</u> 2 °C):	3.6°C
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Actions

Results will be qualified based on the criteria of the following Table:

Table 1. Holding Time Actions for Semivolatile Analyses

	1	ing Time Actions for Semive	Action		
Matrix	Preserved	Criteria	Detected Associated Compounds	Non-Detected Associated Compounds	
	No	≤7 days (for extraction) ≤40 days (for analysis)	Use professi	onal judgment	
	No	> 7 days (for extraction) > 40 days (for analysis)	Ĵ	Use professional judgment	
Aqueous	Yes	≤ 7 days (for extraction) ≤ 40 days (for analysis)	No qualification		
!	Yes	> 7 days (for extraction) > 40 days (for analysis)	.I.	υJ	
	Yes/No	Grossly Exceeded	J	UJ or R	
	No	≤ 14 days (for extraction) ≤ 40 days (for analysis)	Use profession	onal judgment	
Non-Aqueous	No	> 14 days (for extraction) > 40 days (for analysis)	J	Use professional judgment	
	Yes	≤ 14 days (for extraction) ≤ 40 days (for analysis)	No qualification		
	Yes	> 14 days (for extraction) > 40 days (for analysis)	J	UJ	
	Yes/No	Grossly Exceeded	J	UJ or R	

All criteria were met _	X_
Criteria were not met see below	

GC/MS TUNING

The assessment of the tuning results is to determine if the sample instrumentation is within the standard tuning QC limits

- _X__ The DFTPP performance results were reviewed and found to be within the specified criteria.
- _X__ DFTPP tuning was performed for every 12 hours of sample analysis.

If no, use professional judgment to determine whether the associated data should be accepted, qualified or rejected.

Notes: These requirements do not apply when samples are analyzed by the Selected Ion Monitoring (SIM) technique.

All mass spectrometer conditions must be identical to those used during the sample analysis. Background subtraction actions resulting in spectral distortion are unacceptable

Notes: No data should be qualified based of DFTPP failure.

The requirement to analyze the instrument performance check solution is optional when analysis of PAHs/pentachlorophenol is to be performed by the SIM technique.

List	the	samples	affected	

Actions:

- 1. If sample are analyzed without a preceding valid instrument performance check or are analyzed 12 hours after the Instrument Performance Check, qualify all data in those samples as unusable (R).
- 2. If ion abundance criteria are not met, use professional judgment to determine to what extent the data may be utilized.
- 3. State in the Data Review Narrative, decisions to use analytical data associated with DFTPP instrument performance checks not meeting the contract requirements.
- 4. Use professional judgment to determine if associated data should be qualified based on the spectrum of the mass calibration compounds.

All criteria were metX	
Criteria were not met	
and/or see below	

INITIAL CALIBRATION VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

Date of initial calibration:		
Instrument ID numbers:		The state of the s
Matrix/Level:Aqueo	us/iow	
Date of initial calibration:		
Instrument ID numbers:		
Matrix/Level:Aqueo	us/low	
Date of initial calibration:	02/24/16;	_03/02/16_(Scan)
Instrument ID numbers:	GCMSP	· · · · · · · · · · · · · · · · · · ·
Matrix/Level:Aqueo	us/low	

DATE	LAB	FILE	CRITERIA OUT	COMPOUND	SAMPLES
	ID#		RFs, %RSD, %D, r		AFFECTED
			Initial calibration m	eets the required criteria.	

Actions:

Qualify the initial calibration analytes listed in Table 2 using the following criteria:

Table 3. Initial Calibration Actions for Semivolatile Analysis

Criteria	Action		
Criteria	Detect	Non-detect	
Initial Calibration not performed at specified frequency and sequence	Use professional judgment R	Use professional judgment R	
Initial Calibration not performed at the specified concentrations	J	ÜJ	
RRF < Minimum RRF in Table 2 for target analyte	Use professional judgment J+ or R	R	
RRF ≥ Minimum RRF in Table 2 for target analyte	No qualification	No qualification	
%RSD > Maximum %RSD in Table 2 for target analyte	1.	Use professional judgment	
%RSD ≤ Maximum %RSD in Table 2 for target analyte	No qualification	No qualification	

Initial Calibration

Table 2. RRF, %RSD, and %D Acceptance Criteria in Initial Calibration and CCV for Semivolatile Analysis

Analyte	Minimum RRF	Maximum "%RSD	Opening Maximum %D ¹	Opening Maximum %D ¹
1,4-Dioxane	0.010	40.0	-40.0	= 50.0
Benzaldehyde	0.100	40,0	± 40.0	± 50.0
Phenol	0.080	20.0	- 20.0	± 25.0
Bis(2-chloroethyl)ether	0.100	20.0	± 20.0	±25.0
2-Chlorophenol	0.200	20.0	-20.0	±25.0
2-Methylphenol	0.010	20.0	= 20,0	±25.0
3-Methylphenol	0.010	20.0	± 20.0	±25.0
2,2'-Oxybis-(1-chloropropane)	0.010	20.0	±25.0	± 50.0
Acetophenone	0.060	20.0	±20,0	±25.0
4-Methylphenol	0.010	20.0	±20.0	± 25.0
N-Nitroso-di-n-propylamine	0.080	20.0	±25.0	= 25.0
Hexachloroethane	0.100	20.0	= 20.0	- 25.0
Nitrobenzene	0.090	20.0	± 20.0	= 25.0
Isophorone	0.100	20.0	= 20.0	= 25.0
2-Nitrophenol	0.060	20.0	=20.0	= 25.0
2,4-Dimethylphenol	0.050	20.0	±25.0	= 50.0
Bis(2-chloroethoxy)methane	0.080	20.0	± 20.0	= 25.0
2,4-Dichlorophenol	0.060	20,0	±20.0	=25.0
Naphthalene	0.200	20.0	± 20.0	= 25.0
4-Chloroaniline	0.010	40.0	-40.0	= 50.0
lexachlorobutadiene	0.040	20,0	± 20.0	- 25.0
Caprolactam	0.010	40.0	= 30.0	= 50.0
4-Chloro-3-methylphenol	0.040	20,0	± 20.0	= 25.0
2-Methylnaphthalene	0.100	20.0	= 20.0	= 25.0
lexachlorocyclopentadiene	0.010	40.0	= 40.0	= 50.0
2,4,6-Trichlorophenol	0.090	20.0	= 20.0	= 25.0
2,4,5-Trichlorophenol	0.100	20.0	= 20.0	= 25.0
I, I'-Biphenyl	0.200	20.0	= 20.0	= 25.0

Analyte	Minimum RRF	Maximum %RSD	Opening Maximum %D ¹	Opening Maximum %D ¹
2-Chloronaphthalene	0.300	20.0	± 20.0	±25.0
2-Nitroaniline	0.060	20.0	±25.0	±25.0
Dimethylphthalate	0.300	20.0	± 25.0	±25.0
2,6-Dinitrotoluene	0.080	20.0	±20.0	± 25.0
Acenaphthylene	0.400	20.0	±20.0	± 25.0
3-Nitroaniline	0.010	20.0	±25.0	±50.0
Acenaphthene	0.200	20.0	± 20.0	±25.0
2,4-Dinitrophenol	0.010	40.0	±50.0	±50.0
4-Nitrophenol	0.010	40,0	±40.0	± 50.0
Dibenzofuran	0.300	20.0	± 20.0	± 25.0
2,4-Dinitrotoluene	0.070	20.0	=20.0	±25.0
Diethylphthalate	0.300	20.0	= 20.0	±25.0
1,2,4,5-Tetrachlorobenzene	0.100	20.0	±20.0	± 25.0
4-Chlorophenyl-phenylether	0.100	20.0	= 20.0	±25.0
Fluorene	0.200	20.0	± 20.0	±25.0
4-Nitroaniline	0.010	40.0	±40.0	± 50.0
4,6-Dinitro-2-methylphenol	0.010	40.0	±30.0	± 50.0
4-Bromophenyl-phenyl ether	0.070	20.0	±20.0	±25.0
N-Nitrosodiphenylamine	0.100	20.0	± 20.0	± 25.0
Hexachlorobenzene	0.050	20.0	± 20.0	± 25.0
Atrazine	0.010	40.0	± 25.0	±50.0
Pentachlorophenol	0.010	40.0	+40.0	± 50.0
Phenanthrene	0.200	20.0	= 20.0	±25.0
Anthracene	0.200	20.0	± 20.0	±25.0
Carbazole	0.050	20,0	± 20.0	± 25.0
Di-n-butylphthalate	0.500	20.0	=20.0	± 25.0
Fluoranthene	0.100	20.0	± 20.0	±25.0
Pyrene	0.400	20.0	± 25.0	±50.0
Butylbenzylphthalate	0.100	20.0	= 25.0	±50.0

Analyte	Minimum RRF	Maximum %RSD	Opening Maximum %D ¹	Opening Maximum %D ¹
3,3'-Dichlorobenzidine	0.010	40.0	± 40.0	- 50.0
Benzo(a)anthracene	0.300	20.0	± 20.0	± 25.0
Chrysene	0.200	20.0	± 20.0	±50.0
Bis(2-ethylhexyl) phthalate	0.200	20.0	± 25.0	=50.0
Di-n-octylphthalate	0.010	40.0	= 40.0	= 50.0
Benzo(b)fluoranthene	0.010	20.0	±25.0	± 50.0
Benzo(k)fluoranthene	0.010	20,0	= 25.0	= 50.0
Benzo(a)pyrene	0.010	20.0	=20.0	±50.0
Indeno(1,2,3-cd)pyrene	0.010	20.0	± 25.0	± 50.0
Dibenzo(a,h)anthracene	0.010	20.0	± 25.0	± 50.0
Benzo(g,h,i)perylene	0.010	20.0	≟ 30.0	± 50.0
2,3,4,6-Tetrachlorophenol	0.040	20.0	= 20.0	= 50.0
Naphthalene	0.600	20.0	±25.0	± 25.0
2-Methylnaphthalene	0.300	20.0	= 20.0	- 25.0
Acenaphthylene	0.900	20.0	±20.0	= 25.0
Acenaphthene	0.500	20.0	± 20.0	± 25.0
Fluorene	0.700	20.0	±25.0	± 50.0
Phenanthrene	0.300	20.0	= 25.0	± 50.0
Anthracene	0.400	20.0	± 25,0	± 50.0
Fluoranthene	0.400	20.0	± 25.0	± 50.0
Pyrene	0.500	20.0	±30.0	± 50.0
Benzo(a)anthracene	0.400	20.0	= 25.0	= 50.0
Chyrsene	0.400	20,0	= 25.0	= 50.0
Benzo(b)fluoranthene	0.100	20.0	± 30.0	± 50.0
Benzo(k)fluoranthene	0.100	20.0	= 30.0	± 50.0
Benzo(a)pyrene	0.100	20.0	= 25.0	= 50.0
Indeno(1,2,3-cd)pyrene	0.100	20.0	± 40.0	= 50.0
Dibenzo(a,h)anthracene	0.010	25.0	± 40.0	± 50.0
Benzo(g,h,i)perylene	0.020	25.0	± 40.0	± 50.0

Pentachlorophenol	0.010	40.0	= 50.0	± 50.0	
Deuterated Monitoring Compounds					

Analyte	Minimum RRF	Maximum %RSD	Opening Maximum %D ¹	Closing Maximum %D
1,4-Dioxane-d ₈	0.010	20.0	±25.0	± 50.0
Phenol-d₅	0.010	20.0	= 25.0	±25.0
Bis-(2-chloroethyl)ether-d ₈	0.100	20.0	= 20.0	±25.0
2-Chlorophenol-d ₄	0.200	20.0	± 20.0	±25.0
4-Methylphenol-d ₈	0.010	20.0	= 20.0	±25.0
4-Chloroaniline-d ₄	0.010	40.0	= 40.0	± 50.0
Nitrobenzene-d5	0.050	20.0	= 20.0	±25.0
2-Nitrophenol-d4	0.050	20.0	± 20.0	± 25.0
2,4-Dichlorophenol-d;	0.060	20.0	± 20.0	± 25.0
Dimethylphthalate-d ₆	0.300	20.0	= 20.0	± 25.0
Acenaphthylene-d ₈	0.400	20.0	+ 20.0	± 25.0
4-Nitrophenol-d ₁	0.010	40.0	± 40.0	± 50.0
Fluorene-d ₁₀	0,100	20.0	= 20.0	± 25.0
4,6-Dinitro-2-methylphenol-d2	0.010	40.0	- 30.0	± 50.0
Anthracene-d ₁₀	0.300	20.0	± 20.0	± 25.0
Pyrene-d ₁₀	0,300	20.0	-25.0	±50.0
Benzo(a)pyrene-d ₁₂	0.010	20.0	= 20.0	± 50.0
Fluoranthene-d _{in} (SIM)	0.400	20.0	=25.0	±50.0
2-Methylnaphthalene-d ₁₀ (SIM)	0.300	20.0	±20.0	±25.0

If a closing CCV is acting as an opening CCV, all target analytes must meet the requirements for an opening CCV.

Note: If analysis by SIM technique is requested for PAH/pentachlorophenols, calibration standards analyzed at 0.10, 0.20, 0.40, 0.80, and 1.0 ng/uL for each target compound of interest and the associated DMCs. Pentachlorophenol will require only a four point initial calibration at 0.20, 0.40, 0.80, and 1.0 ng/uL.

All criteria were met	X
Criteria were not met	
and/or see below	

CONTINUING CALIBRATION VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

Date of initial calibration:02/01/16_(SIM)	02/24/16;_03/02/16_(Scan)_
Date of initial calibration verification (CCV):_02/01/16	
Date of continuing calibration verification (CCV):_03/16/16	03/16/16
Date of closing CCV:	-
Instrument ID numbers:GCMS4P	GCMSP
Matrix/Level:Aqueous/low	
Date of initial calibration:01/30/16;_02/01/16_(SIM)	
Date of initial calibration verification (CCV):_01/30/16;_02/01/16	
Date of continuing calibration verification (CCV):_03/15/16	
Date of closing CCV:	
Instrument ID numbers:GCMSF	
Matrix/Level:Aqueous/low	

DATE	LAB FILE	CRITERIA OUT	COMPOUND	SAMPLES
	ID#	RFs, %RSD, <u>%D</u> , r		AFFECTED
MSF				
03/15/16	cc6483-25	-20.9	Caprolactam	
		-22.4	2-Nitroaniline	
		-45.1	2,4-Dinitrophenol	
		-29.3	4,6-Dinitro-o-cresol	
MSP				
03/16/15	cc4524-50	-20.3	Caprolactam	
		-21.4	2-Nitroaniline	
	cc4514-50	20.6	Benzaldehyde	
03/17/16	cc4524-25	-25.6	Caprolactam	
		-20.4	2-Nitroaniline	

Note: Continuing calibration verifications %D outside the method criteria but within the guidance document %D required criteria. No closing calibration verification included in data package. No action taken, professional judgment

Actions:

Notes: Verify that the CCV is run at the required frequency (an opening and closing CCV must be run within 12-hour period).

All DMCs must meet the RRF values given in Table 2. No qualification of the data is necessary on DMCs RRF and %RSD/%D alone. Use professional judgment to evaluate DMCs and %RSD/%D data in conjunction with DMCs recoveries to determine the need for qualification of the data.

Qualify the initial calibration analytes listed in Table 2 using the following criteria in the CCVs:

Table 4. CCV Actions for Semivolatile Analysis

Cuitaria for Ononing CCV	Cuitouia fou Clasius CCV	Action		
Criteria for Opening CCV	Criteria for Closing CCV -	Detect	Non-detect	
CCV not performed at required frequency and sequence	CCV not performed at required frequency	Use professional judgment R	Use professional judgment R	
CCV not performed at specified concentration	CCV not performed at specified concentration	Use professional judgment	Use professional judgment	
RRF < Minimum RRF in Table 2 for target analyte	RRF < Minimum RRF in Table 2 for target analyte	Use professional judgment J or R	R	
RRF ≥ Minimum RRF in Table 2 for target analyte	RRF≥ Minimum RRF in Table 2 for target analyte	No qualification	No qualification	
%D outside the Opening Maximum %D limits in Table 2 for target analyte	%D outside the Closing Maximum %D limits in Table 2 for target analyte	J	ŧij	
%D within the inclusive Opening Maximum %D limits in Table 2 for target analyte	%D within the inclusive Closing Maximum %D limits in Table 2 for target analyte	No qualification	No qualification	

All criteria were met_	_X_	_
Criteria were not met		
and/or see below		

CONCENTRATION

BLANK ANALYSIS RESULTS (Sections 1 & 2)

LARID

LEVELL

The assessment of the blank analysis results is to determine the existence and magnitude of contamination problems. The criteria for evaluation of blanks apply only to blanks associated with the samples, including trip, equipment, and laboratory blanks. If problems with any blanks exist, all data associated with the case must be carefully evaluated to determine whether or not there is an inherent variability in the data for the case, or if the problem is an isolated occurrence not affecting other data.

List the contamination in the blanks below. High and low levels blanks must be treated separately.

Notes: The concentration of non-target compounds in all blanks must be less than or equal to 10 ug/L.

The concentration of target compounds in all blanks must be less than its CRQL listed in the method.

Samples taken from a drinking water tap do not have and associated field blank.

COMPOUND

Laboratory blanks

DATE

ANALYZED	LAD ID	MATRIX	COMP COND	UNITS
_No_target_ana			anks.	
Field/Equipmen	t/Trip blank			
DATE Analyzed	LABID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS
			is_data_packageNo_tar	rget_analyte_detected_in
_03/15/16	JC16038-9	_Aqueous/low_	_bis(2-Ethylhexyl)phthala	nte1.2_ug/L

All criteria were met _	_X
Criteria were not met	
and/or see below	_

BLANK ANALYSIS RESULTS (Section 3)

Blank Actions

Qualify samples based on the criteria summarized in Table 5:

Table 5. Blank and TCLP/SPLP LEB Actions for Semivolatile Analysis

Blank Type	Blank Result	Sample Result	Action
	Detect	Non-detect	No qualification
	< CRQL	< CRQL	Report at CRQL and qualify as non-detect (U)
		≥ CRQL	Use professional judgment
	≥ CRQL	< CRQL	Report at CRQL and qualify as non-detect (U)
Method,		≥ CRQL but < Blank Result	Report at sample results and qualify as non-detect (U) or as unusable (R)
TCLP/SPLP LEB, Field		≥ CRQL and ≥ Blank Result	Use professional judgment
	Grossly high	Detect	Report at sample results and qualify as unusable (R)
	TIC > 5.0 ug/L (water) or 0.0050 mg/L (TCLP leachate) or TIC > 170 ug/Kg (soil)	Detect	Use professional judgment

List samples qualified

CONTAMINATION SOURCE/LEVEL	COMPOUND	CONC/UNITS	AL/UNITS	SQL	AFFECTED SAMPLES

All criteria were met _X
Criteria were not met
and/or see below

SURROGATE SPIKE RECOVERIES - DEUTERATED MONITORING COMPOUNDS (DMCs)

Laboratory performance of individual samples is established by evaluation of surrogate spike recoveries – deuterated monitoring compounds. All samples are spiked with surrogate compounds prior to sample analysis. The accuracy of the analysis is measured by the surrogate percent recovery. Since the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the validation of data is frequently subjective and demands analytical experience and professional judgment.

Notes: Recoveries for DMCs in samples and blanks must be within the limits specified in Table 6.

The recovery limits for any of the compounds listed in Table 6 may be expanded at any time during the period of performance if USEPA determines that the limits are too restrictive.

If a DMC is not added in the samples and blanks or the concentrations of DMCs in the samples and blank not the specified, use professional judgment in qualifying the data.

Table 7. DMC Actions for Semivolatile Analysis

	Action		
Criteria	Detect	Non-detect	
%R < 10% (excluding DMCs with 10% as a lower acceptance limit)	J-	R	
10% ≤ %R (excluding DMCs with 10% as a lower acceptance limit) < Lower Acceptance Limit	J-	บม	
Lower Acceptance limit $\leq \%R \leq Upper$ Acceptance Limit	No qualification	No qualification	
%R > Upper Acceptance Limit	J÷	No qualification	

Note: % recovery for Phenol-d5 outside the laboratory control limits but within the guidance document required criteria.

Table 8. Semivolatile DMCs and the Associated Target Analytes

1,4-Dioxane-da (DMC-1)	Phenol-d ₅ (DMC-2)	Bis(2-Chloroethyl) ether-d ₈ (DMC-3)
1,4-Dioxane	Benzaldehyde	Bis(2-chloroethyl)ether
	Phenol	2,2'-Oxybis(1-chloropropane)
		Bis(2-chloroethoxy)methane
2-Chlorophenol-d4(DMC-4)	4-Methylphenol-ds (DMC-5)	4-Chloroaniline-d ₄ (DMC-6)
2-Chlorophenol	2-Methylphenol	4-Chloroaniline
	3-Methylphenol	Hexachlorocyclopentadiene
	4-Methylphenol	Dichlorobenzidine
	2,4-Dimethylphenol	, see
Nitrobenzene-d ₅ (DMC-7)	2-Nitrophenol-d4 (DMC-8)	2,4-Dichlorophenol-d3(DMC-9)
Acetophenone	Isophorone	2,4-Dichlorophenot
N-Nitroso-di-n-propylamine	2-Nitrophenol	Hexachlorobutadiene
Hexachloroethane		Hexachlorocyclopentadiene
Nitrobenzene		4-Chloro-3-methylphenol
2,6-Dinitrotoluene		2,4,6-Trichlorophenol
2,4-Dinitrotoluene		2,4,5-Trichlorophenol
N-Nitrosodiphenylamine		1,2,4,5-Tetrachlorobenzene
3.00		*Pentachlorophenol
		2,3,4,6-Tetrachlorophenol
Dimethylphthalate-d _b (DMC-10)	Acenaphthylene-da (DMC-11)	4-Nitrophenol-d ₄ (DMC-12)
Caprolactam	*Naphthalene	2-Nitroaniline
1,1'-Biphenyl	*2-Methylnaphthalene	3-Nitroaniline
Dimethylphthalate	2-Chloronaphthalene	2,4-Dinitrophenol
Diethylphthalate	*Acenaphthylene	4-Nitrophenol
Di-n-hutylphthalate	*Acenaphthene	4-Nitroaniline
Butylbenzylphthalate		
Bis(2-ethylhexyl) phthalate		
Di-n-octylphthalate		

Fluorene-d ₁₀ (DMC-13)	4,6-Dinitro-2-methylphenol-d ₂ (DMC-14)	Anthracene-d ₁₀ (DMC-15)
Dibenzofuran *Fluorene 4-Chlorophenyl-phenylether 4-Bromophenyl-phenylether Carbazole	4,6-Dinitro-2-methylphenol	Hexachlorobenzene Atrazine *Phenanthrene *Anthracene
Pyrene-d ₁₀ (DMC-16)	Benzo(a)pyrene-d ₁₂ (DMC-17)	
*Fluoranthene *Pyrene *Benzo(a)anthracene *Chrysene	3,3'-Dichlorobenzidine *Benzo(b)fluoranthene *Benzo(k)fluoranthene *Benzo(a)pyrene *Indeno(1,2,3-cd)pyrene *Dibenzo(a,h)anthracene *Benzo(g,h,i)perylene	

^{*}Included in optional Target Analyte List (TAL) of PAHs and PCP only.

Table 9. Semivolatile SIM DMCs and the Associated Target Analytes

Fluoranthene-d10 (DMC-1)	2-Methylnaphthalene-d10 (DMC-2)
Fluoranthene	Naphthalene
Pyrene	2-Methylnaphthalene
Benzo(a)anthracene	Acenaphthylene
Chrysene	Acenaphthene
Benzo(b)fluoranthene	Fluorene
Benzo(k)fluoranthene	Pentachlorophenol
Benzo(a)pyrene	Phenanthrene
Indeno(1,2,3-ed)pyrene	Anthracene
Dibenzo(a,h)anthracene	
Benzo(g,h,i)perylene	

All criteria were met _X	
Criteria were not met	
and/or see below	

VII. A MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD)

This data is generated to determine long term precision and accuracy in the analytical method for various matrices. This data alone cannot be used to evaluate the precision and accuracy of individual samples. If any % R in the MS or MSD falls outside the designated range, the reviewer should determine if there are matrix effects, i.e. LCS data are within the QC limits but MS/MSD data are outside QC limit.

1. MS/MSD Recoveries and Precision Criteria

The laboratory should use one MS and a duplicate analysis of an unspiked field sample if target analytes are expected in the sample. If target analytes are not expected, MS/MSD should be analyzed.

NOTES:

Data for MS and MSDs will not be present unless requested by the

Region.

Notify the Contract Laboratory COR if a field or trip blank was used for the

MS and MSD.

For a Matrix Spike that does not meet criteria, apply the action to only the field sample used to prepare the Matrix Spike sample. If it is clearly stated in the data validation materials that the samples were taken through incremental sampling or some other method guaranteeing the homogeneity of the sample group, then the entire sample group may be qualified.

List the %Rs, RPD of the compounds which do not meet the criteria.

Sample ID:JC16090-1 Sample ID:JC16038-2_(SIM)				Matrix/Level:_Groundwa Matrix/Level:_Groundwa		
MS OR MSD JC16038-2 (SIM)	COMPOUND	% R	RPD	QC LIMITS	ACTION	
_MS/MSD	1,4-dioxane	37%/-70	%	20160	No_action	
Moto: M	o oottom kiek lees	1	1 0 4			

Note: No action, high level of sample relative to amount spiked.

* QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper limit.

* If QC limits are not available, use limits of 70 – 130 %.

Actions:

QUALITY	%R < LL	%R > UL
Positive results	J	J
Nondetects results	R	Accept

MS/MSD criteria apply only to the unspiked sample, its dilutions, and the associated MS/MSD samples:

If the % R for the affected compounds were < LL (or 70 %), qualify positive results (J) and nondetects (UJ).

If the % R for the affected compounds were > UL (or 130 %), only qualify positive results (J).

If 25 % or more of all MS/MSD %R were < LL (or 70 %) or if two or more MS/MSD %Rs were < 10%, qualify all positive results (J) and reject nondetects (R).

A separate worksheet should be used for each MS/MSD pair.

All criteria were met _	Х
Criteria were not met	
and/or see below	-

INTERNAL STANDARD PERFORMANCE

The assessment of the internal standard (IS) parameter is used to assist the data reviewer in determining the condition of the analytical instrumentation.

List the internal standard area of samples which do not meet the criteria.

DATE SAMPLE ID IS OUT IS AREA ACCEPTABLE ACTION RANGE

Internal standard area counts meet the required criteria.

Action:

- 1. If an internal standard area count for a sample or blank is greater than 200.0% of the area for the associated standard (opening CCV or mid-point standard from initial calibration) (see Table 10 below):
 - Qualify detects for compounds quantitated using that internal standard as estimated low (J-).
 - b. Do not qualify non-detected associated compounds.
- 2. If an internal standard area count for a sample or blank is less than 20.0% of the area for the associated standard (opening CCV or mid-point standard from initial calibration):
 - a. Qualify detects for compounds quantitated using that internal standard as estimated high (J+).
 - Qualify non-detected associated compounds as unusable (R).
- If an internal standard area count for a sample or blank is greater than or equal to 50.0%, and less than or equal to 200% of the area for the associated standard opening CCV or mid-point standard from initial calibration, no qualification of the data is necessary.
- 4. If an internal standard RT varies by more than 10.0 seconds: Examine the chromatographic profile for that sample to determine if any false positives or negatives exist. For shifts of a large magnitude, the reviewer may consider partial or total rejection of the data for that sample fraction. Detects should not need to be qualified as unusable (R) if the mass spectral criteria are met.
- If an internal standard RT varies by less than or equal to 10.0 seconds, no qualification of the data is necessary.

Note: Inform the Contract Laboratory Program Project Officer (CLP PO) if the internal standard performance criteria are grossly exceeded. Note in the Data Review Narrative potential effects on the data resulting from unacceptable internal standard performance.

State in the Data Review Narrative if the required internal standard compounds are not added to a sample or blank or if the required internal standard compound is not analyzed at the specified concentration.

Actions:

Table 10. Internal Standard Actions for Semivolatile Analysis

Criteria	Action		
Стега	Detect	Non-detect	
Area response < 20% of the opening CCV or mid-point standard CS3 from ICAL	J+	R	
20% ≤ Area response < 50% of the opening CCV or mid-point standard CS3 from ICAL	J+	ij	
50% ≤ Area response ≤ 200% of the opening CCV or mid-point standard CS3 from ICAL	No qualification	No qualification	
Area response > 200% of the opening CCV or mid-point standard CS3 from ICAL	J-	No qualification	
RT shift between sample/blank and opening CCV or mid-point standard CS3 from ICAL > 10.0 seconds	R	R	
RT shift between sample/blank and opening CCV or mid-point standard CS3 from ICAL < 10.0 seconds	No qualification	No qualification	

		All criteria were metX Criteria were not met and/or see below
TARGET CO	MPOUND IDENTIFICATION	
Criteria:		
	ve Retention Times (RRTs) of reported cor T [opening Continuing Calibration Verificatio ion].	
List compoun	nds not meeting the criteria described above:	
Sample ID	Compounds	Actions
spectrum from	a of the sample compound and a current labor the associated calibration standard (openinust match according to the following criteria: All ions present in the standard mass spectromast because in the sample spectromast of these ions in standard and sample spectra (e.g., for a standard spectrum, the corresponding standard spectrum, the corresponding standard spectrum, must be evaluated.	ng CCV or mid-point standard from initial actrum at a relative intensity greater than rum. The standard from initial actrum at a relative intensity greater than rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a relative intensity greater than a rum. The standard from initial actrum at a rum.
List compoun	spectral interpretation.	,
Sample ID	Compounds	Actions
Sample ID	Compounds ———————————————————————————————————	Actions

Action:

- 1. The application of qualitative criteria for GC/MS analysis of target compounds requires professional judgment. It is up to the reviewer's discretion to obtain additional information from the laboratory. If it is determined that incorrect identifications were made, qualify all such data as unusable (R).
- Use professional judgment to qualify the data if it is determined that cross-contamination has occurred.
- 3. Note in the Data Review Narrative any changes made to the reported compounds or concerns regarding target compound identifications. Note, for Contract Laboratory COR action, the necessity for numerous or significant changes.

TENTATIVELY IDENTIFIED COMPOUNDS (TICS)

NOTE: Tentatively identified compounds should only be evaluated when requested by a party from outside of the Hazardous Waste Support Section (HWSS).

L	.ISt	T	ICs

Sample ID Compound		Sample ID	Compound
	=======================================		=======================================
2000 2000 2000 2000 2000 2000 2000 200			

Action:

- 1. Qualify all TIC results for which there is presumptive evidence of a match (e.g. greater than or equal to 85% match) as tentatively identified (NJ), with approximated concentrations. TICs labeled "unknown" are qualified as estimated (J).
- 2. General actions related to the review of TIC results are as follows:
 - a. If it is determined that a tentative identification of a non-target compound is unacceptable, change the tentative identification to "unknown" or another appropriate identification, and qualify the result as estimated (J).
 - b. If all contractually-required peaks were not library searched and quantitated, the Region's designated representative may request these data from the laboratory.
- In deciding whether a library search result for a TIC represents a reasonable identification, use professional judgment. If there is more than one possible match, report the result as "either compound X or compound Y". If there is a lack of isomer specificity, change the TIC result to a nonspecific isomer result (e.g., 1,3,5-trimethyl benzene to trimethyl benzene isomer) or to a compound class (e.g., 2-methyl, 3-ethyl benzene to a substituted aromatic compound).
- 4. The reviewer may elect to report all similar compounds as a total (e.g., all alkanes may be summarized and reported as total hydrocarbons).

- 5. Target compounds from other fractions and suspected laboratory contaminants should be marked as "non-reportable".
- 6. Other Case factors may influence TIC judgments. If a sample TIC match is poor, but other samples have a TIC with a valid library match, similar RRT, and the same ions, infer identification information from the other sample TIC results.
- 7. Note in the Data Review Narrative any changes made to the reported data or any concerns regarding TIC identifications.
- 8. Note, for Contract Laboratory COR action, failure to properly evaluate and report TICs

All criteria were met _	_X_	
Criteria were not met		
and/or see below		_

SAMPLE QUANTITATION AND REPORTED CONTRACT REQUIRED QUANTITATION LIMITS (CRQLS)

Action:

- 1. When a sample is analyzed at more than one dilution, the lower CRQL are used unless a QC exceedance dictates the use of higher CRQLs from the diluted sample. Samples reported with an "E" qualifier should be reported from the diluted sample.
- 2. If any discrepancies are found, the Region's designated representative may contact the laboratory to obtain additional information that could resolve any differences. If a discrepancy remains unresolved, the reviewer must use professional judgment to decide which value is the most accurate. Under these circumstances, the reviewer may determine that qualification of data is warranted. Note in the Data Review Narrative a description of the reasons for data qualification and the qualification that is applied to the data.
- 3. For non-aqueous samples, if the solids is less than 10.0%, use professional judgment for both detects and non-detects. If the percent solid for a soil sample is greater than or equal to 10.0% and less than 30.0%, use professional judgment to qualify detects and non-detects. If the percent solid for a soil sample is greater than or equal to 30.0%, detects and non-detects should not be qualified (see Table 11).
- 4. Note, for Contract Laboratory COR action, numerous or significant failures to accurately quantify the target compounds or to properly evaluate and adjust CRQLs.
- 5. Results between MDL and CRQL should be qualified as estimated "J".
- 6. Results < MDL should be reported at the CRQL and qualified "U". MDLs themselves should not be reported.

Table 11. Percent Solids Actions for Semivolatile Analysis for Non-Aqueous Samples

Criteria	Action			
Criteria	Detects	Non-detects		
%Solids < 10.0% 10.0%	lac professional judgment lac professional judgment the professional judgment No qualification	lise professional judgment sie professional judgment ie professional judgment No qualification		

SAMPLE QUANTITATION

The sample quantitation evaluation is to verify laboratory quantitation results. In the space below, please show a minimum of one sample calculation:

Sample ID:	JC16038-	1 Analyte:1,4-Dioxane	RF:_0.711_
[]	=	(112855)(40)/(115395)(0.711) 55.0 ppm Ok	

QUANTITATION LIMITS

A. Dilution performed

SAMPLE ID	DILUTION FACTOR	REASON FOR DILUTION
JC16038-8	40 x	1,4-Dioxane outside calibration range
JC16038-10	5 x	1,4-Dioxane outside calibration range

All criteria were met
Criteria were not met
and/or see belowX

FIELD DUPLICATE PRECISION

Sample IDs:JC16038-5/-6 I	Matrix:	Groundwater
---------------------------	---------	-------------

Field duplicates samples may be taken and analyzed as an indication of overall precision. These analyses measure both field and lab precision; therefore, the results may have more variability than laboratory duplicates which only laboratory performance. It is also expected that soil duplicate results will have a greater variance than water matrices due to difficulties associated with collecting identical field duplicate samples.

The project QAPP should be reviewed for project-specific information.

Suggested criteria: if large RPD (> 50 %) is observed, confirm identification of the samples and note differences. If both samples and duplicate are <5 SQL, the RPD criteria is doubled.

COMPOUND	SQL ug/L	SAMPLE CONC.	DUPLICATE CONC.	RPD	ACTION
Acenaphthene	0.29	ND	0.44	-	No action
Fluorene	0.31	ND	0.49		No action
2-Methyl naphthalene	0.30	ND	3.4	-	Qualify the results as estimated (J) in
Naphthalene	0.30	ND	2.3	-	JC16038-5/-6.
Phenanthrene	0.24	ND	0.61		No action

Note: No action taken, professional judgment. Sample and duplicate < 5 SQL.

		All criteria were metX Criteria were not met and/or see below
OTHER ISSUES		
A. System Perfo	rmance	
List samples qualified	based on the degradation of system	performance during simple analysis:
Sample ID	Comments	Actions
Action:		
degraded during sam		etermined that system performance has aboratory Program COR any action as a antly affected the data.
B. Overall Asses	sment of Data	
List samples qualified	based on other issues:	
Sample ID	Comments	Actions
_No_other_issues_th _used_for_decission_		e_dataResults_are_valid_and_can_be

Action:

- 1. Use professional judgment to determine if there is any need to qualify data which were not qualified based on the Quality Control (QC) criteria previously discussed.
- 2. Write a brief narrative to give the user an indication of the analytical limitations of the data. Inform the Contract Laboratory COR the action, any inconsistency of the data with the Sample Delivery Group (SDG) Narrative. If sufficient information on the intended use and required quality of the data is available, the reviewer should include their assessment of the usability of the data within the given context. This may be used as part of a formal Data Quality Assessment (DQA).
- 3. Sometimes, due to dilutions, re-analysis or SIM/Scan runs are being performed, there will be multiple results for a single analyte from a single sample. The following criteria and professional judgment are used to determine which result should be reported:
 - The analysis with the lower CRQL
 - The analysis with the better QC results
 - The analysis with the higher results

EXECUTIVE NARRATIVE

SDG No:

JC16038

Laboratory:

Accutest, New Jersey

Analysis:

SW846-8015C (DAI)

Number of Samples:

10

Location:

BMSMC, Building 5 Area

Humacao, PR

SUMMARY:

Eight (8) groundwater samples, one field blank, and one trip blank were analyzed for the low molecular weight alcohols (LMWAs) list following method SW846-8015C. The sample results were assessed according to USEPA data validation guidance documents in the following order of precedence: "Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW-846 (Final Update III, December 1996)," specifically for Methods 8000/8015C are utilized. The QC criteria and data validation actions listed on the data review worksheets are from the primary guidance document, unless otherwise noted.

Results are valid and can be used for decision making purposes.

Critical issues:

None

Major:

None

Minor:

1. Initial calibration did not meet the method specific criteria for n-butyl alcohol (initial calibration) in column #2. Results reported are from column #1. Isopropanol continuing calibration verification the outside method specific

criteria in one of the columns. No action taken, professional judgment.

Critical findings:

None

Major findings:

None

Minor findings:

None

COMMENTS:

Results are valid and can be used for decision making purposes.

Reviewers Name:

Rafael Infante

Chemist License 1828

Signature:

April 16, 2016

Date:

SAMPLE ORGANIC DATA SAMPLE SUMMARY

Sample ID: JC16038-1

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	-	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	•	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
Methanol	200	ug/l	1.0	-	U	Yes

Sample ID: JC16038-2

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	-	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
Methanoi	200	ug/l	1.0	-	U	Yes

Sample ID: JC16038-3

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	-	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	-	υ	Yes
Methanol	200	uø/l	1.0	_	t)	Ves

Sample ID: JC16038-4

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	-	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
Methanoi	200	ug/l	1.0	_	U	Yes

Sample ID: JC16038-5

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016

Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	-	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	υ	Yes
n-Propyl Alcohol	100	ug/l	1.0	2	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	5	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	2	U	Yes
Methanol	200	ug/l	1.0	7.0	U	Yes

Sammelle Grand Strain Sammelle Grand Strain Sammelle Grand Strain Strain

Sample location: BMSMC Building 5 Area

Sampling Hote: 83/19/2016

Analyte Name	Result	Units D	ilution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	2	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	7.	U	Yes
n-Butyl Alcohol	100	ug/l	1.0		U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	7.1	υ	Yes
Methanol	200	ug/l	1.0		U	Yes

Sample ID: JC16038-7

Sample location: BMSMC Building 5 Area

Sampling date: 3/9/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	2	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	Α.	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	2	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	5	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	2	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	31	U	Yes
Methanol	200	ug/l	1.0	2	U	Yes

Sample ID: JC16038-8

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	-	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
Methanol	200	ug/l	1.0	-	U	Yes

Sample ID: JC16038-9

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	ilution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	-	U	Yes
Isobutyl Alcohol	100	ug/l	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Propyl Alcohol	100	ug/l	1.0	-	U	Yes
n-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
Methanol	200	ug/l	1.0	-	U	Yes

Sample ID: JC16038-10

Sample location: BMSMC Building 5 Area

Sampling date: 3/8/2016 Matrix: Groundwater

METHOD: 8015C

Analyte Name	Result	Units Di	lution Factor	Lab Flag	Validation	Reportable
Ethanol	100	ug/l	1.0	2	U	Yes
Isobutyl Alcohol	100	ug/!	1.0	-	U	Yes
Isopropyl Alcohol	100	ug/l	1.0	2	U	Yes
n-Propyl Alcohol	100	ug/l	1.0		U	Yes
n-Butyl Alcohol	100	ug/l	1.0	-	U	Yes
sec-Butyl Alcohol	100	ug/l	1.0		U	Yes
Methanol	200	ug/l	1.0	2	U	Yes

	Project Number:JC16038 Date:03/08-09/2016
	Shipping Date:03/09-10/2016
	EPA Region:2
REVIEW OF VOLATILE ORG The following guidelines for evaluating volatile organics volations. This document will assist the reviewer in using pelecision and in better serving the needs of the data users. The JSEPA data validation guidance documents in the follow evaluating Solid Waste, Physical/Chemical Methods SV pecifically for Methods 8000/8015C are utilized. The QC collatar eview worksheets are from the primary guidance document in the hardcopied (laboratory name)Accutesteviewed and the quality control and performance data sun	were created to delineate required validation rofessional judgment to make more informed the sample results were assessed according to ing order of precedence: "Test Methods for V-846 (Final Update III, December 1996)," riteria and data validation actions listed on the nent, unless otherwise noted. data package received has been
ncluded:	
ab. Project/SDG No.:JC16038 No. of Samples:10	Sample matrix:Groundwater
rin blank No. IC16028 7	
rip blank No.:JC16038-7 ield blank No.:JC16038-9	
Equipment blank No.:	
ield duplicate No.:JC16038-5/-6_(MW-16/MW-16E	0)
	•
X Data Completeness X Holding Times	X Laboratory Control Spikes
N/A_ GC/MS Tuning	X Field Duplicates X Calibrations
N/A_ Octobs Tulling N/A_ Internal Standard Performance	X Compound Identifications
X Blanks	X Compound Quantitation
X Surrogate Recoveries	X Quantitation Limits
X Surrogate RecoveriesX Matrix Spike/Matrix Spike Duplicate	^ Quantitation Limits
Overall Comments:_Low_molecular_weight_alcohols_t	oy_SW-846_8015C_(DAI)
Definition of Qualifiers:	
- Estimated results	
J- Compound not detected	
R- Rejected data	
J- Estimated nordetect	
Reviewer: 10 4000 angular angu	
GIG. TWILL IN THE IN	

DATA COMPLETENESS

MISSING INFORMATION	DATE LAB. CONTACTED	DATE RECEIVED
100	-	
		
		<u> </u>
-015-040		100

All criteria were met _	_X_	
Criteria were not met		
and/or see below		

HOLDING TIMES

The objective of this parameter is to ascertain the validity of the results based on the holding time of the sample from time of collection to the time of analysis.

Complete table for all samples and note the analysis and/or preservation not within criteria

SAMPLE ID	DATE SAMPLED	DATE ANALYZED	рН	ACTION
	All samples analyzed w	ithin the recommended	method	holding time.
				2082
<u> </u>				
			. [

Criteria

Aqueous samples – 14 days from sample collection for preserved samples (pH \leq 2, 4°C), no air bubbles.

Aqueous samples – 7 days from sample collection for unpreserved samples, 4°C, no air bubbles. Soil samples- 7 days from sample collection.

Cooler temperature (Criteria: 4 + 2 °C): 3.6°C

Actions

If the VOCs vial(s) have air bubbles, estimate positive results (J) and reject nondetects (R).

If the % solids of soil samples is 10-50%, estimates positive results (J) and nondetects (UJ)

If the % solid of soil samples is < 10%, estimate positive results (J) and reject nondetects (R).

If holding times are exceeded but < 14 days beyond criteria, estimate positive results (J) and nondetects (UJ).

If holding times are exceeded but < 28 days beyond criteria, estimate positive results (J) and reject nondetects (R).

If holding times are grossly exceeded (> 28 days beyond criteria), reject all results (R).

If samples were not iced or if the ice were melted (> 10°C), estimate positive results (J) and nondetects (UJ).

		Criteria w	All criteria were metN/A ere not met see below			
GC/MS TUNING						
The assessment o standard tuning QC		o determine if the sample instrumen	tation is within the			
N/A_ The BFB p	erformance results were	e reviewed and found to be within the	specified criteria.			
N/A_ BFB tuning was performed for every 12 hours of sample analysis.						
If no, use profession qualified or rejected		nine whether the associated data sh	nould be accepted,			
List	the	samples	affected:			

If mass calibration is in error, all associated data are rejected.

All criteria were mel	
Criteria were not met	
and/or see belowX	

CALIBRATION VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

Date of initial calibration:	02/29/16	
Dates of continuing calibration	n:_02/29/16 (initial);_03/17/16;_03/18/16	
Instrument ID number:	GCGH	
Matrix/Level:	Aqueous/low	

DATE	LAB FILE ID#	CRITERIA OUT	COMPOUND	SAMPLES
	<u></u>	RFs, %RSD, %D, r		AFFECTED
02/29/16	GH103541.D	26.96% (RSD) (#2)	n-Butanol	-
03/17/16	CC5193-5000	-21.5% (D) (#2)	Isobutanol	-
03/17/16	CC5193-5000	-28.8% (D) (#1)	Isobutanol	-

Note: Initial and continuing calibration meets method specific criteria except for n-butyl alcohol (initial calibration) in column #2. Results reported are from column #1. Isopropanol continuing calibration verification outside method specific criteria in one of the columns. No action taken, professional judgment.

Criteria

All RFs must be > 0.05 regardless of method requirements for SPCC.

All %RSD must be ≤ 15 % regardless of method requirements for CCC.

All %Ds must be ≤ 20% regardless of method requirements for CCC.

It should be noted that Region 2 SOP HW-24 does not specify criterion for the curve correlation coefficient (r). A limit for r of > 0.995 has therefore been utilized as professional judgment.

Actions

If any compound has an initial RF or a continuing RF of < 0.05, estimate positive results (J) and reject nondetects (R), regardless of method requirements.

If any compound has a %RSD > 15%, estimate positive results (J) and use professional judgment to qualify nondetects.

If any compound has a %RSD > 90%, estimate positive results (J) and reject nondetects (R).

If any compound has a % D > 20%, estimate positive results (J) and reject nondetects (R).

If any compound has a % D > 20%, estimate positive results (J) and nondetects (UJ).

If any compound has a % D > 90%, estimate positive results (J) and reject nondetects (R).

If any compound has r < 0.995, estimate positive results and nondetects.

A separate worksheet should be filled for each initial curve

All criteria were met _	X
Criteria were not met	
and/or see below	

V A. BLANK ANALYSIS RESULTS (Sections 1 & 2)

The assessment of the blank analysis results is to determine the existence and magnitude of contamination problems. The criteria for evaluation of blanks apply only to blanks associated with the samples, including trip, equipment, and laboratory blanks. If problems with any blanks exist, all data associated with the case must be carefully evaluated to determine whether or not there is an inherent variability in the data for the case, or if the problem is an isolated occurrence not affecting other data.

List the contamination in the blanks below. High and low levels blanks must be treated separately.

Laboratory blanks

DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS
DATE ANALYZED	LAB ID	LEVEL/ MATRIX	COMPOUND	CONCENTRATION UNITS
	_package			uipment_blank_analyzed
V +4 ·			<u> </u>	

All criteria were met _X
Criteria were not met
and/or see below

VB. BLANK ANALYSIS RESULTS (Section 3)

Blank Actions

Action Levels (ALs) should be based upon the highest concentration of contaminant determined in any blank. Do not qualify any blank with another blank. The ALs for samples which have been diluted should be corrected for the sample dilution factor and/or % moisture, where applicable. No positive sample results should be reported unless the concentration of the compound in the samples exceeds the ALs:

ALs = 10x the amount of common contaminants (methylene chloride, acetone, 2-butanone, and toluene)

ALs = 5x for any other compounds

Specific actions are as follows:

If the concentration is < sample quantitation limit (SQL) and \le AL, report the compound as not detected (U) at the SQL.

If the concentration is \geq SQL but \leq AL, report the compound as not detected (U) at the reported concentration.

If the concentration is \geq SQL and > AL, report the concentration unqualified.

Notes:

High and low level blanks must be treated separately

Compounds qualified "U" for blank contamination are still considered "hits" when qualifying for calibration criteria.

CONTAMINATION SOURCE/LEVEL	COMPOUND	CONC/UNITS	AL/UNITS	SQL	AFFECTED SAMPLES
				_	
			L		

All criteria were metX
Criteria were not met
and/or see below

SURROGATE SPIKE RECOVERIES

Laboratory performance of individual samples is established by evaluation of surrogate spike recoveries. All samples are spiked with surrogate compounds prior to sample analysis. The accuracy of the analysis is measured by the surrogate percent recovery. Since the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the validation of data is frequently subjective and demands analytical experience and professional judgment.

List the percent recoveries (%Rs) which do not meet the criteria for surrogate recovery.

Matrix: solid/aqueous

SAMPLE ID		SURROGAT	E COMPOUND		ACTION
Hex	kanol l	OBFM .	TOL-d8	BFB	
_All_surrogate_recover	ies_within_lat	oratory_cont	rol_limits		442-27-277
QC Limits* (Aqueous)	56 to 145	to	to	to	
QC Limits* (Solid-Low)LL_to_UL	to				
QC Limits* (Solid-Med)LL_to_UL		to	to	to	
1,2-DCA = 1,2-Dichloro DBFM = Dibromofluoro				: Toluene-d8 omofluorobenzen	ne

- * QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper limit.
- * If QC limits are not available, use limits of 80 120 % for aqueous and 70 130 % for solid samples.

Actions:

QUALITY	%R < 10%	%R = 10% - LL	%R > UL
Positive results	J	J	J
Nondetects results	R	UJ	Accept

Surrogate action should be applied:

If one or more surrogate in the VOC fraction is out of specification, but has a recovery of > 10%. If any one surrogate in a fraction shows < 10 % recovery.

All criteria were met _X
Criteria were not met
and/or see below

VII. A MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD)

This data is generated to determine long term precision and accuracy in the analytical method for various matrices. This data alone cannot be used to evaluate the precision and accuracy of individual samples. If any % R in the MS or MSD falls outside the designated range, the reviewer should determine if there are matrix effects, i.e. LCS data are within the QC limits but MS/MSD data are outside QC limit.

MS/MSD Recoveries and Precision Criteria

The laboratory should use one MS and a duplicate analysis of an unspiked field sample if target analytes are expected in the sample. If target analytes are not expected, MS/MSD should be analyzed.

List the %Rs, RPD of the compounds which do not meet the criteria.

	5796-1MS/-1MSD_ 6038-1MS/-1MSD_				Groundwater Groundwater	
MS OR MSD	COMPOUND	% R	RPD	QC LIMITS	ACTION	
MS/MSD%_red	coveries_and_RPD_	within_lab	oratory_	control_limits		
			-			

- * QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper limit.
- * If QC limits are not available, use limits of 70 130 %.

Actions:

QUALITY	%R < LL	%R > UL
Positive results	J	J
Nondetects results	R	Accept

MS/MSD criteria apply only to the unspiked sample, its dilutions, and the associated MS/MSD samples:

If the % R for the affected compounds were < LL (or 70 %), qualify positive results (J) and nondetects (UJ).

If the % R for the affected compounds were > UL (or 130 %), only qualify positive results (J).

If 25 % or more of all MS/MSD %R were < LL (or 70 %) or if two or more MS/MSD %Rs were < 10%, qualify all positive results (J) and reject nondetects (R).

A separate worksheet should be used for each MS/MSD pair.

All criteria were met _	х_
Criteria were not met	
and/or see below	39

VII. B MATRIX SPIKE/MATRIX SPIKE DUPLICATE

MS/MSD - Unspiked Compounds

It should be noted that Region 2 SOP HW-24 does not specify a MS/MSD criteria for the unspiked compounds in the sample. A %RSD of < 50% has therefore been utilized as professional judgment.

If all target analytes were spiked in the MS/MSD, this review element is not applicable.

List the %RSD of the compounds which do not meet the criteria.

Sample ID:			Matrix/Level/Unit:		
COMPOUND	SAMPLE CONC.	MS CONC.	MSD CONC.	% RSD	ACTION
-					2 20 0000

Actions:

^{*} If the % RSD > 50, qualify the positive result in the unspiked samples as estimated (J).

^{*} If the % RSD is not calculated (NC) due to nondetected value, use professional judgment to qualify the data.

All criteria were met_	X_
Criteria were not met	
and/or see below	

VIII. LABORATORY CONTROL SAMPLE (LCS) ANALYSIS

This data is generated to determine accuracy of the analytical method for various matrices.

1. LCS Recoveries Criteria

Where LCS spiked with the same analyte at the same concentrations as the MS/MSD? **Yes** or No. If no make note in data review memo.

List the %R of compounds which do not meet the criteria

	LCS ID	COMPOUND	% R	QC LIMH	
Recoveries_within_laboratory_control_limits					
			<u> </u>		
				5	

- * QC limits are laboratory in-house performance criteria, LL = lower limit, UL = upper limit.
- * If QC limits are not available, use limits of 70 130 %.

Actions:

QUALITY	%R < LL	%R > UL
Positive results	J	J
Nondetects results	R	Accept

All analytes in the associated sample results are qualified for the following criteria.

If 25 % of the LCS recoveries were < LL (or 70 %), qualify all positive results (j) and reject nondetects (R).

If two or more LCS were below 10 %, qualify all positive results as (J) and reject nondetects (R).

2. Frequency Criteria:

Where LCS analyzed at the required frequency and for each matrix? <u>Yes</u> or No. If no, the data may be affected. Use professional judgment to determine the severity of the effect and qualify data accordingly. Discuss any actions below and list the samples affected.

				All criteria were metX Criteria were not mel and/or see below
IX.	FIELD/LABOR	ATORY DUPLICATE PRECISION		
	Sample IDs:	_JC16038-5/-6	Matrix:	Groundwater

Field/laboratory duplicates samples may be taken and analyzed as an indication of overall precision. These analyses measure both field and lab precision; therefore, the results may have more variability than laboratory duplicates which only laboratory performance. It is also expected that soil duplicate results will have a greater variance than water matrices due to difficulties associated with collecting identical field duplicate samples.

The project QAPP should be reviewed for project-specific information. Suggested criteria: RPD \pm 30% for aqueous samples, RPD \pm 50 % for solid samples. If both samples and duplicate are <5 SQL, the RPD criteria is doubled.

COMPOUND	SQL	SAMPLE CONC.	DUPLICATE CONC.	RPD	ACTION	
RPD within laboratory and generally acceptable control limits.						

Actions:

Qualify as estimated positive results (J) and nondetects (UJ) for the compound that exceeded the above criteria. For organics, only the sample and duplicate will be qualified.

If an RPD cannot be calculated because one or both of the sample results is not detected, the following actions apply:

If one sample result is not detected and the other is greater than 5x the SQL qualify (J/UJ).

If one sample value is not detected and the other is greater than 5x the SQL and the SQLs for the sample and duplicate are significantly different, use professional judgment to determine if qualification is appropriate.

If one sample value is not detected and the other is less than 5x, use professional judgment to determine if qualification is appropriate.

If both sample and duplicate results are not detected, no action is needed.

Actions:

All criteria were metNA	_
Criteria were not met	
and/or see below	

X. INTERNAL STANDARD PERFORMANCE

The assessment of the internal standard (IS) parameter is used to assist the data reviewer in determining the condition of the analytical instrumentation.

List the internal standard area of samples which do not meet the criteria.

- * Area of +100% or -50% of the IS area in the associated calibration standard.
- * Retention time (RT) within 30 seconds of the IS area in the associated calibration standard.

DATE	SAMPLE ID	IS OUT	IS AREA	ACCEPTABLE ACTION RANGE
	_			

1. IS actions should be applied to the compound quantitated with the out-of-control ISs

QUALITY	IS AREA < -25%	IS AREA = -25 % TO - 50%	IS AREA > + 100%
Positive results	J	J	J
Nondetected results	R	UJ	ACCEPT

2. If a IS retention time varies more than 30 seconds, the chromatographic profile for that sample must be examined to determine if any false positive or negative exists. For shifts of a large magnitude, the reviewer may consider partial or total rejection of the data for the sample fraction.

All criteria were met _	X_
Criteria were not met	
and/or see below	

XII. SAMPLE QUANTITATION

The sample quantitation evaluation is to verify laboratory quantitation results. In the space below, please show a minimum of one sample calculation:

JC16038-1

Hexanol

RF = 127.5

[] = (477585)/(127.5)

= 3745.8 ppb OK

All criteria were met _X
Criteria were not met
and/or see below

XII. QUANTITATION LIMIT	S
-------------------------	---

A. Dilution performed

SAMPLE ID	DILUTION FACTOR	REASON FOR DILUTION		

-	
В.	Percent Solids
	List samples which have ≤ 50 % solids
Action	
	If the % solids of a soil sample is 10-50%, estimate positive results (J) and nondetects (U.
	If the % solids of a soil sample is < 10%, estimate positive results (J) and reject nondetection (R)

EXECUTIVE NARRATIVE

SDG No:

JC16038

Laboratory:

Accutest, New Jersey

Analysis:

SW846-8081B

Number of Samples:

Location:

BMSMC, Building 5 Area

Humacao, PR

SUMMARY:

Five (5) groundwater samples were analyzed for selected pesticides following method SW846-8081B. The sample results were assessed according to USEPA data validation guidance documents in the following order of precedence Hazardous Waste Support Section SOP No. HW-36A, Revision O, June, 2015. SOM02.2. Pesticide Data Validation. The QC criteria and data validation actions listed on the data review worksheets are

from the primary guidance document, unless otherwise noted.

Results are valid and can be used for decision making purposes.

Critical issues:

None

Major:

None

Minor:

1. No MS/MSD duplicate analyzed with this data package. LCS/LCSD used to

assess accuracy; % recoveries and RPD within laboratory control limits.

Critical findings:

None

Major findings:

None

Minor findings:

None

COMMENTS:

Results are valid and can be used for decision making purposes.

Reviewers Name:

Rafael Infante

Chemist License 1888

Signature:

April 16, 2016

Date:

SAMPLE ORGANIC DATA SAMPLE SUMMARY

Sample ID: JC16038-8

Sample location: BMSMC Building 5 Area

Sampling date: 8-Mar-16
Matrix: Groundwater

METHOD: 8081B

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
beta-BHC	0.010	ug/l	1	-	-	Yes
4,4'-DDD	0.010	ug/l	1	-	-	Yes
4,4'-DDT	0.010	ug/l	1	-	-	Yes

Sample ID: JC16038-9

Sample location: BMSMC Building 5 Area

Sampling date: 8-Mar-16 Matrix: Groundwater

METHOD: 8081B

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
beta-BHC	0.010	ug/l	1	-	U	Yes
4,4'-DDD	0.010	ug/l	1	-	U	Yes
4,4'-DDT	0.010	ug/l	1	-	U	Yes

Sample ID: JC16038-10

Sample location: BMSMC Building 5 Area

Sampling date: 8-Mar-16
Matrix: Groundwater

METHOD: 8081B

Analyte Name	Result	Units	Dilution Factor	Lab Flag	Validation	Reportable
beta-BHC	0.011	ug/l	1	-	-	Yes
4,4'-DDD	0.011	ug/l	1	-	-	Yes
4,4'-DDT	0.011	ug/l	1	_	-	Yes

	Project/Case Number:JC16038 Sampling Date:March_8,_2016
	Shipping Date:March_10,_2016
	EPA Region No.:22
REVIEW OF PESTICI	DE ORGANIC PACKAGE
required validation actions. This document judgment to make more informed decision users. The sample results were assessed a documents in the following order of precede https://doi.org/10.100/1001/1001/1001/1001/1001/1001/1	volatile organics were created to delineate will assist the reviewer in using professional and in better serving the needs of the data according to USEPA data validation guidance ence Hazardous Waste Support Section SOP No. Pesticide Data Validation. The QC criteria and a review worksheets are from the primary d.
The hardcopied (laboratory name) _Accutest reviewed and the quality control and performance da	data package received has been ta summarized. The data review for VOCs included:
Lab. Project/SDG No.:JC16038 No. of Samples:3	
Trip blank No.:	
X Data CompletenessX Holding TimesN/A GC/MS TuningX Internal Standard PerformanceX BlanksX Surrogate RecoveriesX Matrix Spike/Matrix Spike Duplicate	X Laboratory Control SpikesX Field DuplicatesX CalibrationsX Compound IdentificationsX Compound QuantitationX Quantitation Limits
Overall Comments:Selected_pesticides_by_S	W846-8081B
Definition of Qualifiers: J- Estimated results U- Compound not detected R- Rejected data UJ- Estimated nondetect	

Reviewer:__ Rafuel Defaut _____ Date:__April_16,_2016______

DATA COMPLETENESS

MISSING INFORMATION	DATE LAB. CONTACTED	DATE RECEIVED
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	The second secon	
		<u> </u>

All criteria were met _X
Criteria were not met
and/or see below

HOLDING TIMES

The objective of this parameter is to ascertain the validity of the results based on the holding time of the sample from time of collection to the time of analysis.

Complete table for all samples and note the analysis and/or preservation not within criteria

SAMPLE ID	DATE SAMPLED	DATE EXTRACTED/ANALYZED	ACTION

Preservatives:	_All_samples	_extracted_and	_analyzed_within_	_the_required_c	criteria

Criteria

Aqueous samples - seven (7) days from sample collection for extraction; 40 days from sample collection for analysis.

Non-aqueous samples – fourteen (14) days from sample collection for extraction; 40 days from sample collection for analysis.

Cooler temperature (Criteria: 4 ± 2 °C): 3.6°C - OK

Actions

Qualify aqueous sample results using preservation and technical holding time information as follows:

- a. If there is no evidence that the samples were properly preserved (T = 4° C \pm 2° C), and the samples were extracted or analyzed within the technical holding times, qualify detects as estimated (J) and non-detects as estimated (UJ).
- b. If there is no evidence that the samples were properly preserved (T = 4° C \pm 2° C), and the samples were extracted or analyzed outside the technical holding times, qualify detects as estimated (UJ) and non-detects as estimated (UJ).
- c. If the samples were properly preserved, and were extracted and analyzed within the technical holding times, no qualification of the data is necessary.
- d. If the samples were properly preserved, and were extracted or analyzed outside the technical holding times, qualify detects as estimated (J) and non-detects as estimated (UJ). Note in the Data Review Narrative that holding times were exceeded and the effect of exceeding the holding time on the resulting data.

- e. Use professional judgment to qualify samples whose temperature upon receipt at the laboratory is either below 2 degrees centigrade or above 6 degrees centigrade.
- f. If technical holding times are grossly exceeded, use professional judgment to qualify the data.

Qualify non-aqueous sample results using preservation and technical holding time information as follows:

- a. If there is no evidence that the samples were properly preserved (T = 4° C \pm 2° C), and the samples were extracted or analyzed within the technical holding time, qualify detects as estimated (J) and non-detects as estimated (UJ).
- b. If there is no evidence that the samples were properly preserved ($T = 4^{\circ}C \pm 2^{\circ}C$), and the samples were extracted or analyzed outside the technical holding time, qualify detects as estimated (UJ).
- c. If the samples were properly preserved, and were extracted and analyzed within the technical holding time, no qualification of the data is necessary.
- d. If the samples were properly preserved, and were extracted or analyzed outside the technical holding time, qualify detects as estimated (J) and non-detects as estimated (UJ). Note in the Data Review Narrative that holding times were exceeded and the effect of exceeding the holding time on the resulting data.
- e. Use professional judgment to qualify samples whose temperature upon receipt at the laboratory is either below 2 degrees centigrade or above 6 degrees centigrade.
- f. If technical holding times are grossly exceeded, use professional judgment to qualify the data.

	All criteria were metX	
Criteria	were not met see below	

GAS CHROMATOGRAPH WITH ELECTRON CAPTURE DETECTOR (GC/ECD) INSTRUMENT PERFORMANCE CHECK (SECTIONS 1 TO 5)

1. Resolution Check Mixture

Criteria

Is the resolution between two adjacent peaks in the Resolution Check Mixture C greater than or equal to 80.0% for all analytes for the primary column and greater than or equal to 50.0% for the confirmation column? Yes? or No?

Is the resolution between two adjacent peaks in the Resolution Check Mixture (A and B) greater than or equal to 60.0%? Yes? or No?

Note: If resolution criteria are not met, the quantitative results may not be accurate due to inadequate resolution. Qualitative identifications may also be questionable if coelution exists.

Action

- a. Qualify detects for target compounds that were not adequately resolved as tentatively identified
- b. Qualify non-detected compounds as unusable (R).

2. Performance Evaluation Mixture (PEM) Resolution Criteria

Criteria

Is PEM analysis performed at the required frequency (at the end of each pesticide initial calibration sequence and every 12 hours)? Yes? or No?

Action

a. If PEM is not performed at the required frequency, qualify all associated sample and blank results as unusable (R).

Criteria

Is PEM % Resolution < 90%?

Yes? or No?

Action

- a. a. Qualify detects for target compounds that were not adequately resolved as tentatively identified (NJ).
- b. Qualify non-detected compounds as unusable (R).

	All criteria were met	.x
Criteria	were not met see below.	

3. PEM 4,4'-DDT Breakdown

Criteria

Is the PEM 4,4'-DDT % Breakdown >20.0% and 4,4'-DDT is detected? Yes? or No?

Action

a. Qualify detects for 4,4'-DDT; detects for 4,4'-DDD; and detects for 4,4'-DDE as estimated (J)

Criteria

Is the PEM 4,4'-DDT % Breakdown >20.0% and 4,4'-DDT is not detected

Yes? or No?

Action

- a. Qualify non-detects for 4,4'- DDT as unusable (R)
- b. Qualify detects for 4,4'-DDD as tentatively identified (NJ)
- c. Qualify detects for 4,4'-DDE as tentatively identified (NJ)

4. PEM Endrin Breakdown

Criteria

Is the PEM Endrin % Breakdown >20.0% and Endrin is detected?

Yes? or No?

Action

a. Qualify detects for Endrin; detects for Endrin aldehyde; and detects for Endrin ketone as estimated (J)

Criteria

Is the PEM Endrin % Breakdown >20.0% and Endrin is not detected

Yes? or No?

Action

- a. Qualify non-detects for Endrin as unusable (R)
- b. Qualify detects for Endrin aldehyde as tentatively identified (NJ)
- c. Qualify detects for Endrin ketone as tentatively identified (NJ)

	All criteria were met	x
Criteria	were not met see below.	2.75

5. Mid-point Individual Standard Mixture Resolution -

Criteria

Is the resolution between two adjacent peaks in the Resolution Check Mixture C greater than or equal to 80.0% for all analytes for the primary column and greater than or equal to 50.0% for the confirmation column?

Yes? or No?

Is the resolution between two adjacent peaks in the Resolution Check Mixture (A and B) greater than or equal to 90.0%?

Yes? or No?

Note: If resolution criteria are not met, the quantitative results may not be accurate due to inadequate resolution. Qualitative identifications may also be questionable if coelution exists.

Action

- a. Qualify detects for target compounds that were not adequately resolved as tentatively identified (NJ).
- b. Qualify non-detected compounds as unusable (R).

Criteria

Is mid-point individual standard mixture analysis performed at the required frequency (every 12 hours)?

Yes? or No?

Action

a. If the mid-point individual standard mixture analysis is not performed at the required frequency, qualify all associated sample and blank results as unusable (R).

All criteria were metX_	_
Criteria were not met	
and/or see below	

CALIBRATION VERIFICATION

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing and maintaining acceptable quantitative data.

	о оор.	2010 01	producing and maina	ining doodplebic qualitative se	aua.
			Date of initial calibration	on:03/18/16	
			Dates of continuing ca	alibration:03/18/16_(initial); rs:GC4G	_03/20/16;_03/21/16_
				Aqueous/low	
				·	
DATE	LAB ID#	FILE	CRITERIA OUT RFs, %RSD, %D, r	COMPOUND	SAMPLES AFFECTED
	ID#		1(1 5, 70((0), 700, 1		AFFECTED
Initial a	nd conti	inuing o		equired criteria. Closing calibra required criteria	tion performed and
Criteria					
Are a five HW-36A, f				h concentration levels as show	n in Table 3 of SOP <u>Yes</u> ? or No?
Actions					
If the stand effect on th		ncentrat	ions listed in Table 3 a	re not used, use professional jud	dgment to evaluate the
Criteria					
Are RT Wi	ndows	calcula	ted correctly?		Yes? or No?
Action					
Recalculat	e the w	indows	and use the corrected	d values for all evaluations.	

Criteria

Are the Percent Relative Standard Deviation (%RSD) of the CFs for each of the single component target compounds less than or equal to 20.0%, except for alpha-BHC and delta-BHC?

Yes? or No?

Are the %RSD of the CFs for alpha-BHC and delta-BHC less than or equal to 25.0%. Yes? or No?

Is the %RSD of the CFs for each of the Toxaphene peaks must be < 30% when 5-point ICAL is performed?

Yes? or No?

Is the %RSD of the CFs for the two surrogates (tetrachloro-m-xylene and decachlorobiphenyl) less than or equal to 30.0%.

Yes? or No?

Action

- a. If the %RSD criteria are not met, qualify detects as estimated (J) and use professional judgment to qualify non-detected target compounds.
- b. If the %RSD criteria are within allowable limits, no qualification of the data is necessary

Continuing Calibration Checks

Criteria

Is the continuing calibration standard analyzed at the acceptable time intervals? Yes? or No?

Action

- a. If more than 14 hours has elapsed from the injection of the instrument blank that begins an analytical sequence (opening CCV) and the injection of either a PEM or mid-point concentration of the Individual Standard Mixtures (A and B) or (C), qualify all data as unusable (R).
- b. If more than 12 hours has elapsed from the injection of the instrument blank that begins an analytical sequence (opening CCV) and the injection of the last sample or blank that is part of the same analytical sequence, qualify all data as unusable (R).
- c. If more than 72 hours has elapsed from the injection of the sample with a Toxaphene detection and the Toxaphene Calibration Verification Standard (CS3), qualify all data as unusable (R).

Criteria

Is the Percent Difference (%D) within ±25.0% for the PEM sample?

Yes? or No?

Action

a. Qualify associated detects as estimated (J) and non-detects as estimated (UJ).

Criteria

For the Calibration Verification Standard (CS3); is the Percent Difference (%D) within ±25.0%?

Yes? or No?

Action

Qualify associated detects as estimated (J) and non-detects as estimated (UJ).

All criteria were metX	
Criteria were not met	
and/or see below	

Criteria

Is the PEM 4,4'-DDT % Breakdown >20.0% and 4,4'-DDT is detected?

Yes? or No?

Action

- a. Qualify detects for 4,4'-DDT; detects for 4,4'-DDD; and detects for 4,4'-DDE as estimated (J)
- b. Non-detected associated compounds are not qualified

Criteria

Is the PEM 4,4'-DDT % Breakdown >20.0% and 4,4'-DDT is not detected

Yes? or No?

Action

- a. Qualify non-detects for 4,4'- DDT as unusable (R)
- b. Qualify detects for 4,4'-DDD as tentatively identified (NJ)
- c. Qualify detects for 4,4'-DDE as tentatively identified (NJ)

Criteria

Is the PEM Endrin % Breakdown >20.0% and Endrin is detected?

Yes? or No?

Action

- a. Qualify detects for Endrin; detects for Endrin aldehyde; and detects for Endrin ketone as estimated (J)
- b. Non-detected associated compounds are not qualified

Criteria

Is the PEM Endrin % Breakdown >20.0% and Endrin is not detected

Yes? or No?

Action

- a. Qualify non-detects for Endrin as unusable (R)
- b. Qualify detects for Endrin aldehyde as tentatively identified (NJ)
- c. Qualify detects for Endrin ketone as tentatively identified (NJ)

A separate worksheet should be filled for each initial curve

All criteria were met _X
Criteria were not met
and/or see below

BLANK ANALYSIS RESULTS (Sections 1 & 2)

The assessment of the blank analysis results is to determine the existence and magnitude of contamination problems. The criteria for evaluation of blanks apply only to blanks associated with the samples, including trip, equipment, and laboratory blanks. If problems with any blanks exist, all data associated with the case must be carefully evaluated to determine whether or not there is an inherent variability in the data for the case, or if the problem is an isolated occurrence not affecting other data.

List the contami	nation in the bla	anks below. Hig	h and low levels blanks	must be treated separately.
CRQL concentra	ationN	/A		
Laboratory blank	ks			
DATE Analyzed	LAB ID	LEVEL! MATRIX	COMPOUND	CONCENTRATION UNITS
			2,000 No. 20 20 10 10 10 10 10 10 10 10 10 10 10 10 10	nit_of_0.01_and_0.001_ug/L.
Field/Equipment	t/Trip blank			
DATE Analyzed	LAB ID	LEVEL/ Matrix	COMPOUND	CONCENTRATION UNITS
_package	<u> </u>			nk_analyzed_with_this_data_
			WW 2	
			_	
				20 C C C C C C C C C C C C C C C C C C C

All criteria were met _X_	
Criteria were not met	
and/or see below	

BLANK ANALYSIS RESULTS (Section 3)

Blank Actions

Action Levels (ALs) should be based upon the highest concentration of contaminant determined in any blank. Do not qualify any blank with another blank. The ALs for samples which have been diluted should be corrected for the sample dilution factor and/or % moisture, where applicable. No positive sample results should be reported unless the concentration of the compound in the samples exceeds the ALs:

The concentration of non-target compounds in all blanks must be less than or equal to 10 μ g/L. The concentration of each target compound found in the method or field blanks must be less than its CRQL listed in the method.

Data concerning the field blanks are not evaluated as part of the CCS process. If field blanks are present, the data reviewer should evaluate this data in a similar fashion as the method blanks.

Specific actions are as follows:

Blank Actions for Pesticide Analyses

Blank Type	Blank Result	Sample Result	Action for Samples	
	Detects	Not detected	No qualification required	
	< CRQL	< CRQL	Report CRQL value with a U	
		≥CRQL	No qualification required	
Method, Sulfur		< CRQL	Report CRQL value with a U	
Cleanup, Instrument, Field, TCLP/SPLP	> CRQL = CRQL	≥ CRQL and ≤ blank concentration	Report blank value for sample concentration with a U	
		≥ CRQL and > blank concentration	No qualification required	
		≤CRQL	Report CRQL value with a U	
		> CRQL	No qualification required	
	Gross contamination	Detects	Report blank value for sample concentration with a U	

All criteria were met	X
Criteria were not met	
and/or see below	

CONTAMINATION SOURCE/LEVEL	COMPOUND	CONC/UNITS	AL/UNITS	SQL	AFFECTED SAMPLES
		_			

All criteria were met __X__ Criteria were not met and/or see below ____

SURROGATE SPIKE RECOVERIES

Laboratory performance of individual samples is established by evaluation of surrogate spike recoveries. All samples are spiked with surrogate compounds prior to sample analysis. The accuracy of the analysis is measured by the surrogate percent recovery. Since the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the validation of data is frequently subjective and demands analytical experience and professional judgment.

List the percent recoveries (%Rs) which do not meet the criteria for surrogate recovery.

Matrix:_Ground	lwater	77. (1)				_
Lab Sample ID	Lab File ID	Sl a	Sl b	S2 a	S2 b	
JC16038-8 JC16038-9 JC16038-10 OP92108-BS1 OP92108-BSD OP92108-MB1	4G66288.D 4G66287.D 4G66289.D 4G66278.D 4G66279.D 4G66277.D	97 78 69 75 86 84	99 86 76 83 97 85	101 67 72 78 94 99	103 73 81 89 103 94	
Surrogate Compounds S1 = Tetrachloro S2 = Decachloro	•	Recove Limits 26-132 10-118	:%			

- (a) Recovery from GC signal #1
- (b) Recovery from GC signal #2

Note: Surrogate recoveries within laboratory control limits.

Actions:

- a. For any surrogate recovery greater than 150%, qualify detected target compounds as biased high (J+).
- b. Do not qualify non-detected target compounds for surrogate recovery > 150 %.
- c. If both surrogate recoveries are greater than or equal to 30% and less than or equal to 150%, no qualification of the data is necessary.
- d. For any surrogate recovery greater than or equal to 10% and less than 30%, qualify detected target compounds as biased low (J-).
- e. For any surrogate recovery greater than or equal to 10% and less than 30%, qualify non-detected target compounds as approximated (UJ).
- f. If low surrogate recoveries are from sample dilution, professional judgment should be used to determine if the resulting data should be qualified. If sample dilution is not a factor:
 - i. Qualify detected target compounds as biased low (J-).
 - ii. Qualify non-detected target compounds as unusable (R).

- g. If surrogate RTs in PEMs, Individual Standard Mixtures, samples, and blanks are outside of the RT Windows, the reviewer must use professional judgment to qualify data.
- h. If surrogate RTs are within RT windows, no qualification of the data is necessary.
- i. If the two surrogates were not added to all samples, MS/MSDs, standards, LCSs, and blanks, use professional judgment in qualifying data as missing surrogate analyte may not directly apply to target analytes.

Summary Surrogate Actions for Pesticide Analyses

	Action*		
Criteria	Detected Target Compounds	Non-detected Target Compounds	
%R > 150%	J+	No qualification	
30% < %R < 150%	No qualification		
10% < %R < 30%	J-	UJ	
%R < 10% (sample dilution not a factor)	J-	R	
%R < 10% (sample dilution is a factor)	Use professional judgment		
RT out of RT window	Use professional judgment		
RT within RT window	No qualification		

^{*} Use professional judgment in qualifying data, as surrogate recovery problems may not directly apply to target analytes.

All criteria were met _N/A____ Criteria were not met and/or see below

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD)

This data is generated to determine long term precision and accuracy in the analytical method for various matrices. This data alone cannot be used to evaluate the precision and accuracy of individual samples. If any % R in the MS or MSD falls outside the designated range, the reviewer should determine if there are matrix effects, i.e. LCS data are within the QC limits but MS/MSD data are outside QC limit.

1. MS/MSD Recoveries and Precision Criteria

List the %Rs, RPD of the compounds which do not meet the criteria.

Data for MS and MSDs will not be present unless requested by the Region.

Notify the Contract Laboratory Program Project Officer (CLP PO) if a field blank was used for the MS and MSD, unless designated as such by the Region.

NOTE: For a Matrix Spike that does not meet criteria, apply the action to only the field sample used to prepare the Matrix Spike sample. If it is clearly stated in the data validation materials that the samples were taken through incremental sampling or some other method guaranteeing the homogeneity of the sample group, then the entire sample group may be qualified.

Sample ID: Matrix/Level:			Level:		
MS OR MSD	COMPOUND	% R	RPD	QC LIMITS	ACTION
					_assess_accuracy;_%_
Action					

No qualification of the data is necessary on MS and MSD data alone. However, using professional judgment, the validator may use the MS and MSD results in conjunction with other QC criteria and determine the need for some qualification of the data.

A separate worksheet should be used for each MS/MSD pair.

All criteria were met _	_X
Criteria were not met	
and/or see below	

LABORATORY CONTROL SAMPLE (LCS) ANALYSIS

This data is generated to determine accuracy of the analytical method for various matrices.

LCS Recoveries Criteria

LCS Spike Compound	Recovery Limits (%)
gamma-BHC	50 – 120
Heptachlor epoxide	50 – 150
Dieldrin	30 – 130
4,4'-DDE	50 – 150
Endrin	50 – 120
Endosulfan sulfate	50 – 120
trans-Chlordane	30 – 130
Tetrachloro-m-xylene (surrogate)	30 – 150
Decachlorobiphenyl (surrogate)	30 – 150

LCS	concentrations:	:0.25_ug/L		
List the %R	of compounds w	hich do not meet the criteria	3	
	LCS ID	COMPOUND	% R	QC LIMIT

Action

The following guidance is suggested for qualifying sample data for which the associated LCS does not meet the required criteria.

- a. If the LCS recovery exceeds the upper acceptance limit, qualify detected target compounds as estimated (J). Do not qualify non-detected target compounds.
- b. If the LCS recovery is less than the lower acceptance limit, qualify detected target compounds as estimated (J) and non-detects as unusable (R).
- c. Use professional judgment to qualify data for compounds other than those compounds that are included in the LCS.
- d. Use professional judgment to qualify non-LCS compounds. Take into account the compound class, compound recovery efficiency, analytical problems associated with each compound, and comparability in the performance of the LCS compound to the non-LCS compound.
- e. If the LCS recovery is within allowable limits, no qualification of the data is necessary.

2. Frequency Criteria:

Where LCS analyzed at the required frequency and for each matrix? <u>Yes</u> or No. If no, the data may be affected. Use professional judgment to determine the severity of the effect and qualify data accordingly. Discuss any actions below and list the samples affected.

All criteria were met
Criteria were not met
and/or see belowN/A

FLORISIL CARTRIDGE PERFORMANCE CHECK

NOTE: Florisil cartridge cleanup is mandatory for all extracts.

Criteria

Is the Florisil cartridge performance check conducted at least once on each lot of cartridges used for sample cleanup or every 6 months, whichever is most frequent?

Yes? or No?

Criteria

Are the results for the Florisil Cartridge Performance Check solution included with the data package?

Yes? or No?

Note: If % criteria are not met, examine the raw data for the presence of polar interferences and use professional judgment in qualifying the data as follows:

Action:

- a. If the Percent Recovery is greater than 120% for any of the pesticide target compounds in the Florisil Cartridge Performance Check, qualify detected compounds as estimated (J). Do not qualify non-detected target compounds.
- b. If the Percent Recovery is greater than or equal to 80% and less than or equal to 120% for all the pesticide target compounds, no qualification of the data is necessary.
- c. If the Percent Recovery is greater than or equal to 10% and less than 80% for any of the pesticide target compounds in the Florisil Cartridge Performance Check, qualify detected target compounds as estimated (J) and non-detected target compounds as approximated (UJ).
- d. If the Percent Recovery is less than 10% for any of the pesticide target compounds in the Florisil Cartridge Performance Check, qualify detected compounds as estimated (J) and qualify non-detected target compounds as unusable (R).
- e. If the Percent Recovery of 2,4,5-trichlorophenol in the Florisil Cartridge Performance Check is greater than or equal to 5%, use professional judgment to qualify detected and non-detected target compounds, considering interference on the sample chromatogram.

Note: State in the Data Review Narrative potential effects on the sample data resulting from the Florisil Cartridge Performance Check analysis not yielding acceptable results.

Note: No information for florisil cartridge performance check included in data package. No qualification of the data performed, professional judgment.

All criteria were met	_
Criteria were not met	
and/or see belowN/A_	

GEL PERMEATION CHROMATOGRAPHY (GPC) PERFORMANCE CHECK

NOTE: GPC cleanup is mandatory for all soil samples.

If GPC criteria are not met, examine the raw data for the presence of high molecular weight contaminants; examine subsequent sample data for unusual peaks; and use professional judgment in qualifying the data. Notify the Contract Laboratory Program Project Officer (CLP PO) if the laboratory chooses to analyze samples under unacceptable GPC criteria.

Action:

- a. If the Percent Recovery is less than 10% for the pesticide compounds and surrogates during the GPC calibration check, the non-detected target compounds may be suspect, qualify detected compounds as estimated (J).
- b. If the Percent Recovery is less than 10% for the pesticide compounds and surrogates during the GPC calibration check, qualify all non-detected target compounds as unusable (R).
- c. If the Percent Recovery is greater than or equal to 10% and is less than 80% for any of the pesticide target compounds in the GPC calibration, qualify detected target compounds as estimated (J) and non-detected target compounds as approximated (UJ).
- d. If the Percent Recovery is greater than or equal to 80% and less than or equal to 120% for all the pesticide target compounds, no qualification of the data is necessary.
- e. If high recoveries (i.e., greater than 120%) were obtained for the pesticides and surrogates during the GPC calibration check, qualify detected compounds as estimated (J). Do not qualify non-detected target compounds.

Note: State in the Data Review Narrative potential effects on the sample data resulting from the GPC cleanup analyses not yielding acceptable results.

Note:_ No information for performance of GPC cleanup included in data package. No qualification of the data performed, professional judgment.

All criteria were met	_X
Criteria were not met	
and/or see below	-

TARGET COMPOUND IDENTIFICATION

Criteria:

- 1. Is Retention Times (RTs) of both of the surrogates and reported target compounds in each sample within the calculated RT Windows on both columns?

 Yes? or No?
- 2. Is the Tetrachloro-m-xylene (TCX) RT ±0.05 minutes of the Mean RT (RT) determined from the initial calibration and Decachlorobiphenyl (DCB) within ±0.10 minutes of the RT determined from the initial calibration?

 Yes? or No?
- 3. Is the Percent Difference (%D) for the detected mean concentrations of a pesticide target compound between the two Gas Chromatograph (GC) columns within the inclusive range of ± 25.0 %?

 Yes? or No?
- 4. When no analytes are identified in a sample; are the chromatograms from the analyses of the sample extract and the low-point standard of the initial calibration associated with those analyses on the same scaling factor?

 Yes? or No?
- 5. Does the chromatograms display the Single Component Pesticides (SCPs) detected in the sample and the largest peak of any multi-component analyte detected in the sample at less than full scale.

 Yes? or No?
- 6. If an extract is diluted; does the chromatogram display SCPs peaks between 10-100% of full scale, and multi-component analytes between 25-100% of full scale?

 Yes? or No?
- 7. For any sample; does the baseline of the chromatogram return to below 50% of full scale before the elution time of alpha-BHC, and also return to below 25% of full scale after the elution time of alpha-BHC and before the elution time of DCB?

 Yes? or No?
- 8. If a chromatogram is replotted electronically to meet these requirements; is the scaling factor used displayed on the chromatogram, and both the initial chromatogram and the replotted chromatogram submitted in the data package.

 Yes? or No?

Action:

- a. If the qualitative criteria for both columns were not met, all target compounds that are reported as detected should be considered non-detected.
- b. Use professional judgment to assign an appropriate quantitation limit using the following guidance:
 - If the detected target compound peak was sufficiently outside the pesticide RT Window, the reported values may be a false positive and should be replaced with the sample Contract Required Quantitation Limits (CRQL) value.

- ii. If the detected target compound peak poses an interference with potential detection of another target peak, the reported value should be considered and qualified as unusable (R).
- c. If the data reviewer identifies a peak in both GC column analyses that falls within the appropriate RT Windows, but was reported as a non-detect, the compound may be a false negative. Use professional judgment to decide if the compound should be included.

Note: State in the Data Review Narrative all conclusions made regarding target compound identification.

- d. If the Toxaphene peak RT windows determined from the calibration overlap with SCPs or chromatographic interferences, use professional judgment to qualify the data.
- e. If target compounds were detected on both GC columns, and the Percent Difference between the two results is greater than 25.0%, consider the potential for coelution and use professional judgment to decide whether a much larger concentration obtained on one column versus the other indicates the presence of an interfering compound. If an interfering compound is indicated, use professional judgment to determine how best to report, and if necessary, qualify the data according to these guidelines.
- f. If Toxaphene exhibits a marginal pattern-matching quality, use professional judgment to establish whether the differences are due to environmental "weathering" (i.e., degradation of the earlier eluting peaks relative to the later eluting peaks). If the presence of Toxaphene is strongly suggested, report results as presumptively present (N).

GAS CHROMATOGRAPH/MASS SPECTROMETER (GC/MS) CONFIRMATION

NOTE: This confirmation is not usually provided by the laboratory. In cases where it is provided, use professional judgment to determine if data qualified with "C" can be salvaged if it was previously qualified as unusable (R).

Action:

- a. If the quantitative criteria for both columns were met (≥ 5.0 ng/µL for SCPs and ≥ 125 ng/µL for Toxaphene), determine whether GC/MS confirmation was performed. If it was performed, qualify the data using the following guidance:
 - i. If GC/MS confirmation was not required because the quantitative criteria for both columns was not met, but it was still performed, use professional judgment when evaluating the data to decide whether the detect should be qualified with "C".
 - ii. If GC/MS confirmation was performed, but unsuccessful for a target compound detected by GC/ECD analysis, qualify those detects as "X".

All criteria were met _	_X
Criteria were not met	
and/or see below	100

COMPOUND QUANTITATION AND REPORTED CONTRACT REQUIRED QUANTITATION LIMITS (CRQLS)

The sample quantitation evaluation is to verify laboratory quantitation results. In the space below, please show a minimum of one sample calculation:

op92108-bs1 (Blank Spike)

4,4'-DDD

RF = 0.764

 $[] = (72050664)(50)/(255.3\times10^6)(0.764)$

= 18.47 ppb Ok

Action:

- a. If sample quantitation is different from the reported value, qualify result as unusable (R).
- b. When a sample is analyzed at more than one dilution, the lowest CRQLs are used unless a QC exceedance dictates the use of the higher CRQLs from the diluted sample.
- c. Replace concentrations that exceed the calibration range in the original analysis by crossing out the "E" and its corresponding value on the original reporting form and substituting the data from the diluted sample.
- d. Results between the MDL and CRQL should be qualified as estimated (J).
- e. Results less than the MDL should be reported at the CRQL and qualified (U). MDLs themselves are not reported.
- f. For non-aqueous samples, if the percent moisture is less than 70.0%, no qualification of the data is necessary. If the percent moisture is greater than or equal to 70.0% and less than 90.0%, qualify detects as estimated (J) and non-detects as approximated (UJ). If the percent moisture is greater than or equal to 90.0%, qualify detects as estimated (J) and non-detects as unusable (R) (see Table).

Percent Moisture Actions for Pesticide Analysis for Non-Aqueous Samples

Criteria	Action		
	Detected Associated Compounds	Non-detected Associated Compounds	
% Moisture < 70.0	No qualification		
70.0 < % Moisture < 90.0	J	UJ	
% Moisture > 90.0	J	R	

amples which have <u>s</u>	50 % solids			
		 		
	-			

Note: If any discrepancies are found, the Region's designated representative may contact the laboratory to obtain additional information that could resolve any differences. If a discrepancy remains unresolved, the reviewer must use professional judgment to decide which value is the most accurate. Under these circumstances, the reviewer may determine that qualification of data is warranted. Note in the Data Review Narrative a description of the reasons for data qualification and the qualification that is applied to the data.

Dilution performed

SAMPLE ID	DILUTION FACTOR	REASON FOR DILUTION
	· ·	

All criteria were met	N/A
Criteria were not met	
and/or see below	

FIELD DUPLICATE PRECISION

NOTE: In the absence of QAPP guidance for validating data from field duplicates, the following action will be taken.

Field duplicates samples may be taken and analyzed as an indication of overall precision. These analyses measure both field and lab precision; therefore, the results may have more variability than laboratory duplicates which only laboratory performance. It is also expected that soil duplicate results will have a greater variance than water matrices due to difficulties associated with collecting identical field duplicate samples. Identify which samples within the data package are field duplicates. Estimate the relative percent difference (RPD) between the values for each compound. If large RPDs (> 50%) is observed, confirm identification of samples and note difference in the executive summary.

Sample ID	s:	<u> </u>	 	Matrix:	
COMPOUND	SQL ug/L	SAMPLE CONC.	DUPLICATE CONC.	RPD	ACTION
No field/laboratory assess precision. F				/LCSD % re	covery RPD used to

Actions:

- a. Qualify as estimated positive results (J) and nondetects (UJ) for the compound that exceeded the above criteria. For organics, only the sample and duplicate will be qualified.
- b. If an RPD cannot be calculated because one or both of the sample results is not detected, the following actions apply:
 - i. If one sample result is not detected and the other is greater than 5x the SQL qualify (J/UJ).
 - ii. If one sample value is not detected and the other is greater than 5x the SQL and the SQLs for the sample and duplicate are significantly different, use professional judgment to determine if qualification is appropriate.
 - iii. If one sample value is not detected and the other is less than 5x, use professional judgment to determine if qualification is appropriate.
 - iv. If both sample and duplicate results are not detected, no action is needed.

OVERALL ASSESSMENT OF DATA

Action:

- 1. Use professional judgment to determine if there is any need to qualify data which were not qualified based on the Quality Control (QC) criteria previously discussed.
- 2. Write a brief narrative to give the user an indication of the analytical limitations of the data.

Note: The Contract Laboratory Program Project Officer (CLP PO) must be informed if any inconsistency of the data with the Sample Delivery Group (SDG) Narrative. If sufficient information on the intended use and required quality of the data is available, the reviewer should include their assessment of the usability of the data within the given context. This may be used as part of a formal Data Quality Assessment (DQA).

Overall assessment of the data: Results are valid; the data can be used for

decision making purposes.